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(54) Title: FUNGICIDE COMPOSITIONS FOR PROTECTING FRUITS

(54) Titre: COMPOSITIONS FONGICIDES POUR LA PROTECTION DES FRUITS

(57) Abstract

The invention concerns fungicide compositions useful for fighting against phytopathogenic fungi infesting or capable of infesting fruits, containing: a) at least a fungicide compound inhibiting mitochondrial respiration; and b) at least a fungicide compound inhibiting sterol biosynthesis. The invention also concerns a method for treating fruits against fungal diseases using said compositions.

(57) Abrégé

Compositions fongicides utiles pour luter contre les champignons phytopathogènes infestant ou susceptibles d'infester les fruits, contenant: a) au moins un composé fongicide inhibiteur de la respiration mitochondriale, et b) au moins un composé fongicide inhibiteur de la biosynthèse des stérols, et procédé de traitement des fruits contre les maladies fongiques au moyen des dites compositions.

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Compositions fongicides pour la protection des fruits

- [001] La présente invention concerne de nouvelles compositions fongicides utiles dans le traitement des fruits contre les maladies fongiques, ainsi que les méthodes de traitement des fruits contre les maladies fongiques au moyen des dites compositions.
- [002] Il est bien connu que les fruits, après qu'ils ont été récoltés, se recouvrent de moisissures, ce qui entraîne leur pourriture.

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- 10 [003] Ces moisissures apparaissent après un temps plus ou moins long, selon la nature du fruit ou du légume et les conditions environnantes, et sont le résultat du développement de un ou plusieurs champignons phytopathogènes. Ces champignons sont pour la plupart bien connus, de même que leur mode de développement.
- 15 [004] Ainsi, de nombreuses méthodes de traitement des fruits sont déjà connues et décrites.
 - [005] Parmi celles-ci, on peut citer l'application par trempage et/ou brossage des fruits après récolte dans de solutions de fongicides connus. On se reportera par exemple aux travaux de A. Chitzanidis et coll., *Bulletin OEPP*, 20(1), (1990), 163-168 ou encore à ceux de E. Cohen et coll., *Phytoparasitica*, 18(1), (1990), 17-26.
 - [006] D'autres procédés recommandent un premier traitement de pulvérisation de composés fongicides sur les arbres fruitiers, puis une application sur les fruits par trempage et/ou brossage d'autres fongicides. Ce type de traitement est notamment décrit par S. Toker et coll., *Turk. J. Agric. For.*, 20(1), (1996), 78-83.
 - [007] Parmi les fongicides les plus couramment utilisés pour le traitement des fruits après récolte, on peut citer l'imazalil (décrit dans "The Pesticide Manual", 10th edition, British Crop Protection Council, page 580), le thiabendazole (*ibid.* page 972) ou encore le SOPP (o-phénylphénate de sodium, *ibid.* page 794), qui se sont avérés jusqu'à présent les plus efficaces.
 - [008] Ces composés sont en effet particulièrement actifs contre les souches de *Penicillium*, qui sont à l'origine de nombreuses maladies fongiques.
 - [009] Malheureusement, des souches de *Penicillium* résistantes au thiabendazole (P. R. Harding Jr., *Plant Dis. Rep.*, 56(3), (1972),256-260), et plus récemment résistantes à
- l'imazalil sont apparues (voir par exemple les publications de J. W. Eckert, Phytopathology, 77(12), (1987), 1728, et ISPP Chemical Control Newsletter, 10, (1988), 36-38).

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[0010] Ces composés sont de plus faiblement actifs, voire inactifs sur d'autres champignons phytopathogènes. En outre, si l'imazalil peut être employé en traitement préventif et curatif, cet effet curatif reste limité dans le temps.

5 [0011] Il est également toujours souhaitable d'améliorer les produits fongicides utilisés pour traiter les fruits.

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[0012] Il est également toujours souhaitable de réduire les doses de produits chimiques appliqués sur les fruits, notamment en réduisant les doses d'application des produits.
[0013] Il est enfin toujours désirable d'augmenter la gamme de produits antifongiques à la disposition afin de trouver parmi ceux-ci les mieux adaptés à des usages spécifiques.

[0014] Un premier objet de la présente invention consiste à fournir des compositions fongicides possédant un large spectre d'activité, c'est-à-dire possédant une activité substantielle sur un nombre de champignons phytopathogènes plus important que le nombre de champignons phytopathogènes traités par les compositions connues.

[0015] Un deuxième objet de l'invention consiste à fournir des compositions fongicides possédant à la fois un effet préventif et un effet curatif, en particulier un effet curatif amélioré par rapport aux compositions fongicides actuellement utilisées.

[0016] Un autre objet de la présente invention consiste à fournir des compositions fongicides qui parviennent à lutter efficacement contre les souches de champignons résistantes aux compositions fongicides connues.

[0017] Un autre but de la présente invention est de fournir des compositions fongicides qui sont efficaces à des doses substantiellement plus faibles par rapport aux doses actuellement appliquées.

[0018] Un autre objet de la présente invention consiste à fournir des compositions fongicides appliquées sur des fruits afin d'empêcher ou de retarder leur pourrissement, tout en restant propres à la consommation.

[0019] D'autres objets de l'invention apparaîtront dans l'exposé de l'invention qui est présenté dans la suite de la présente description.

[0020] De façon surprenante, il a été découvert que l'ensemble de ces objets est atteint en totalité ou partie grâce aux compositions fongicides objet de la présente invention.

[0021] La présente invention concerne donc des compositions fongicides utiles pour lutter contre les champignons phytopathogènes infestant ou susceptibles d'infester les fruits, caractérisée en ce qu'elles contiennent :

- au moins un composé fongicide inhibiteur de la respiration mitochondriale, et
- au moins un composé fongicide inhibiteur de la biosynthèse des stérols.

[0022] Parmi les composés fongicides inhibiteurs de la respiration mitochondriale, on peut citer par exemple la strobilurine et dérivés ou composés analogues, comme par exemple l'azoxystrobine, le krésoxim-méthyl, la trifloxystrobine, la picoxystrobine, la discostrobine, mais aussi le 4-chloro-2-cyano-N,N-dimethyl-5-p-tolylimidazole-1-sulfonamide, la 5-méthyl-5-(4-phénoxyphényl)-3-(phénylamino)-2,4-oxazolidinedione (ou Famoxadone) ainsi que les composés de formule générale (I):

$$(Y)_n$$
 $(Y)_n$
 $(Y)_n$
 $(Y)_n$

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dans laquelle:

- M représente un atome d'oxygène ou de soufre ;
- n est un nombre entier égal à 0 ou 1 ;
- Y est un atome de fluor ou de chlore, ou un radical méthyle.

[0023] Les composés de formule (I) sont connus, notamment par la demande de brevet EP-A-0 629 616.

[0024] De façon préférée, on choisira parmi les composés de formule (I), le composé pour lequel M représente un atome de soufre et n est égal à 0, c'est-à-dire la (4-S)-4-méthyl-2-méthylthio-4-phényl-1-phénylamino-2-imidazoline-5-one, dénommée, Fénamidone.

[0025] La liste des composés fongicides inhibiteurs de la respiration mitochondriale ne doit pas être considérée comme limitative, mais comme illustrant, pour l'homme du métier, la gamme de composés inhibiteurs de respiration mitochondriale dont il dispose.

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[0026] De même, à titre d'illustration, parmi les composés inhibiteurs de la biosynthèse des stérols qui peuvent être utilisés dans les compositions fongicides de la présente invention, on peut citer par exemple, l'imazalil.

- [0027] De façon préférée, on choisira, pour les compositions fongicides objet de l'invention, comme inhibiteur de respiration mitochondriale, un composé de formule (I) ou la famoxadone, et comme inhibiteur de la biosynthèse des stérols, l'imazalil.
- [0028] De manière tout particulièrement préférée, les compositions fongicides selon la présente invention comprennent l'imazalil en association avec la fénamidone.
 - [0029] Les compositions selon la présente invention se présentent sous la forme de liquides plus ou moins visqueux, allant d'une viscosité semblable à celle de l'eau, d'huile, jusqu'à des formulations de type cires.
 - [0030] En règle générale, les doses des composés fongicides présents dans les compositions de l'invention dépendent de la nature des composés eux-mêmes, de la nature des fruits à traiter, ainsi que de la nature des maladies à traiter et de leur degré d'infestation.

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- [0031] Les doses d'inhibiteurs de respiration mitochondriale utilisées pour les compositions fongicides de la présente invention sont avantageusement comprises entre 10 mg/l et 1000 mg/l, de préférence comprises entre 20 mg/l et 300 mg/l, de préférence encore entre 40 mg/l et 150 mg/l, par exemple entre 50 mg/l et 100 mg/l.
- [0032] Les doses d'inhibiteurs de biosynthèse des stérols utilisées pour les compositions fongicides de la présente invention sont avantageusement comprises entre 100 mg/l et 3000 mg/l, de préférence comprises entre 50 mg/l et 2500 mg/l, de préférence encore entre 200 mg/l et 2000 mg/l, par exemple entre environ 400 mg/l et 1000 mg/l.
- [0033] Il est bien entendu que les compositions fongicides selon la présente invention peuvent contenir, outre un ou plusieurs inhibiteurs de la respiration mitochondriale et un ou plusieurs inhibiteurs de la biosynthèse des stérols, un ou plusieurs autres composés fongicides connus de l'homme du métier et adaptés aux traitements des maladies fongiques des fruits.

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[0034] À titre d'exemple non limitatif, les autres composés fongicides qui peuvent être inclus dans les compositions de l'invention, comprennent l'acide phosphoreux, ainsi que ses dérivés et ses sels. Un produit tout particulièrement adapté aux compositions de la présente invention est le sel d'aluminium de l'acide phosphoreux, dénommé Fosétyl-Al (décrit dans "The Pesticide Manual", 10th edition, British Crop Protection Council, page 530).

[0035] Ainsi, une composition tout particulièrement préférée de la présente invention comprend imazalil, fénamidone et fosétyl-Al.

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[0036] Les doses de ces autres composés fongicides qui peuvent être ajoutés aux compositions selon la présente invention sont fonction des types de maladies à traiter, de leur degré d'infestation, de la nature des fruits à traiter et de la nature elle-même de ces composés. L'homme du métier saura apprécier les doses à appliquer qui pourront par exemple être comprises entre 500 mg/l et 6000 mg/l, par exemple entre 2000 mg/l et 4000 mg/l.

[0037] Lorsque le fosétyl-Al est utilisé, la dose d'emploi est généralement comprise entre 1000 mg/l et 1500 mg/l, de préférence environ 1200 mg/l.

[0038] Il a donc été découvert de façon surprenante que l'ajout d'un inhibiteur de la respiration mitochondriale à un inhibiteur de la biosynthèse des stérols augmente substantiellement le spectre d'activité des compositions fongicides et notamment possède une efficacité inattendue sur les souches de champignons phytopathogènes résistantes aux inhibiteurs de la biosynthèse des stérols.

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[0039] Ainsi, les compositions fongicides selon l'invention sont efficaces pour traiter la majeure partie des champignons phytopathogènes infestant ou susceptibles d'infester les fruits et notamment:

- Phytophthora spp., par exemple la pourriture brune des agrumes (Phytophthora parasitica), et la gommose des citrus (Phytophthora citrophthora);
- Penicillium spp., par exemple la moisissure bleue (Penicillium italicum), et la moisissure verte (Penicillium digitatum);
- la pourriture amère des agrumes (Geotrichum candidum) ;
- le black-rot des agrumes (Alternaria citri) ;
- l'anthracnose (Colleotrichum gloeosporioides);
- la mélanose ou pourriture phomopsienne (Diplodia natalensis ou Phomopsis citri).

[0040] D'autres champignons phytopathogènes peuvent également être contrôlés à l'aide des compositions fongicides de l'invention. La nature de ces autres champignons dépend en partie de la nature des autres fongicides présents dans les compositions objet de l'invention.

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[0041] Les fruits qui peuvent être traités par les compositions selon l'invention sont de tout type, et particulièrement ceux qui peuvent être endommagés par l'apparition des champignons phytopathogènes décrits plus haut, notamment lors d'un stockage prolongé des dits fruits.

[0042] De plus les compositions selon l'invention trouvent une utilisation particulièrement avantageuse dans le traitement des fruits après récolte, afin d'empêcher ou de retarder leur pourrissement, tout en restant propres à la consommation.
[0043] Ainsi, parmi les fruits qui peuvent être traités avec les compositions fongicides de l'invention on peut citer notamment les agrumes, par exemple, citrons, oranges,

pamplemousses, cédrats, clémentines, mandarines, etc.

[0044] Il apparaît ainsi que le champ d'application des compositions fongicides décrites ici n'est pas limité, et que ces compositions peuvent être employées dans tous les cas de figure où une protection ou une action contre les attaques fongiques est nécessaire afin d'empêcher ou d'enrayer le pourrissement de fruits comestibles.

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[0045] Les compositions selon l'invention comprennent, outres les composés fongicides décrits auparavant, les supports solides ou liquides, acceptables dans le traitement fongique des fruits et/ou les agents tensioactifs également acceptables dans le traitement fongique des fruits. En particulier sont utilisables les supports inertes et usuels et les agents tensioactifs usuels. Ces compositions recouvrent non seulement les compositions prêtes à être appliquées sur les fruits à traiter par trempage ou au moyen d'un dispositif adapté, mais également les compositions concentrées commerciales qui doivent être diluées avant application sur les fruits.

30 [0046] Ces compositions fongicides selon l'invention peuvent contenir aussi toute sorte d'autres ingrédients tels que, par exemple, des colloïdes protecteurs, des adhésifs, des épaississants, des agents thixotropes, des agents de pénétration, des stabilisants, des séquestrants, des agents de texture, des agents de saveur, des exhausteurs de goût, des sucres, des édulcorants, des colorants, etc... Plus généralement, les matières actives peuvent être combinées à tous les additifs solides ou liquides correspondant aux

peuvent être combinées à tous les additifs solides ou liquides correspondant aux techniques habituelles de la mise en formulation.

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[0047] D'une façon générale, les compositions selon l'invention contiennent habituellement de 0,05 à 95 % (en poids) de matière active (par matière active, on entend ici l'ensemble des composés fongicides compris dans la présente formulation), un ou plusieurs supports solides ou liquides et, éventuellement, un ou plusieurs agents tensioactifs.

[0048] Par le terme "support", dans le présent exposé, on désigne une matière organique ou minérale, naturelle ou synthétique, avec laquelle la matière active est combinée pour faciliter son application sur les fruits. Ce support est donc généralement inerte et il doit être acceptable dans le domaine agro-alimentaire. Le support peut être solide (argiles, silicates naturels ou synthétiques, silice, résines, cires, engrais solides, etc...) ou liquide (eau, alcools, notamment le butanol, etc...).

[0049] L'agent tensioactif peut être un agent émulsionnant, dispersant ou mouillant de type ionique ou non ionique ou un mélange de tels agents tensioactifs. On peut citer par exemple des sels d'acides polyacryliques, des sels d'acides lignosulfoniques, des sels d'acides phénolsulfoniques ou naphtalènesulfoniques, des polycondensats d'oxyde d'éthylène sur des alcools gras ou sur des acides gras ou sur des amines grasses, des phénols substitués (notamment des alkylphénols ou des arylphénols), des sels d'esters d'acides sulfosucciniques, des dérivés de la taurine (notamment des alkyltaurates), des esters phosphoriques d'alcools ou de phénols polyoxyéthylés, des esters d'acides gras et de polyols, les dérivés à fonction sulfates, sulfonates et phosphates des composés précédents. La présence d'au moins un agent tensioactif est généralement indispensable lorsque la matière active et/ou le support inerte ne sont pas solubles dans l'eau et que l'agent vecteur de l'application est l'eau.

[0050] Ainsi donc, les compositions selon l'invention peuvent contenir la matière active dans de très larges limites, allant de 0,05 % à 95 % (en poids). Leur teneur en agent tensioactif est avantageusement comprise entre 0,5 % et 40 % en poids. Sauf indication contraire les pourcentages donnés dans cette description sont des pourcentages pondéraux.

[0051] Ces compositions selon l'invention sont elles-mêmes sous des formes assez diverses, solides ou liquides.

[0052] Comme formes de compositions solides, on peut citer les poudres pour poudrage (à teneur en matière active pouvant aller jusqu'à 100 %) et les granulés, notamment ceux obtenus par extrusion, par compactage, par imprégnation d'un support granulé, par

granulation à partir d'une poudre (la teneur en matière active dans ces granulés étant entre 0,5 et 80 % pour ces derniers cas). De telles compositions solides peuvent éventuellement être mises sous forme liquide plus ou moins visqueux, selon le type d'application désiré, par exemple, par dilution dans l'eau.

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[0053] Comme formes de compositions liquides ou destinées à constituer des compositions liquides lors de l'application, on peut citer les solutions, en particulier les concentrés solubles dans l'eau, les émulsions, les suspensions concentrées, les poudres mouillables (ou poudre à pulvériser), les huiles et les cires.

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[0054] Les suspensions concentrées, applicables en pulvérisation, sont préparées de manière à obtenir un produit fluide stable ne se déposant pas et elles contiennent habituellement de 10 à 75 % de matière active, de 0,5 à 15 % d'agents tensioactifs, de 0,1 à 10 % d'agents thixotropes, de 0 à 10 % d'additifs appropriés, comme des anti-mousses, des inhibiteurs de corrosion, des stabilisants, des agents de pénétration et des adhésifs et, comme support, de l'eau ou un liquide organique dans lequel la matière active est peu ou pas soluble : certaines matières solides organiques ou des sels minéraux peuvent être dissous dans le support pour aider à empêcher la sédimentation ou comme antigels pour l'eau.

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[0055] A titre d'exemple, voici une composition de suspension concentrée : Exemple SC 1 :

• - matière active		500 g
• - phosphate de tristyrylphénol polyéthoxylé		50 g
• - alkylphénol polyéthoxylé	50 g	
•- polycarboxylate de sodium		20 g
• - éthylène glycol		50 g
• - huile organopolysiloxanique (antimousse)		1 g
• - polysaccharide		1,5 g
• - eau	316,5	g

[0056] Les poudres mouillables (ou poudre à pulvériser) sont habituellement préparées de manière qu'elles contiennent 20 à 95 % de matière active, et elles contiennent habituellement, en plus du support solide, de 0 à 30 % d'un agent mouillant, de 3 à 20 % d'un agent dispersant, et, quand c'est nécessaire, de 0,1 à 10 % d'un ou plusieurs stabilisants et/ou autres additifs, comme des agents de pénétration, des adhésifs, ou des agents antimottants, colorants, etc...

[0057] Pour obtenir les poudres à pulvériser ou poudres mouillables, on mélange intimement les matières actives dans les mélangeurs appropriés avec les substances additionnelles et on broie avec des moulins ou autres broyeurs appropriés. On obtient par-là des poudres à pulvériser dont la mouillabilité et la mise en suspension sont avantageuses ; on peut les mettre en suspension avec de l'eau à toute concentration désirée et ces suspensions sont utilisables très avantageusement en particulier pour l'application par exemple sur les feuilles des végétaux ou sur les semences.

10 [0058] A titre d'exemple, voici une composition pour poudre mouillable (ou poudre à pulvériser):

Exemple PM 1

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• - matière active 50%

• - alcool gras éthoxylé (agent mouillant) 2,5%

• - phényléthylphénol éthoxylé (agent dispersant) 5%

•- farine (support inerte) 42,5%

[0059] Les dispersions et émulsions aqueuses, par exemple les compositions obtenues en diluant à l'aide d'eau une poudre mouillable selon l'invention, sont comprises dans le cadre général de la présente invention. Les émulsions peuvent être du type eau-dans-l'huile ou huile-dans-l'eau et elles peuvent avoir une consistance épaisse comme celle d'une "mayonnaise", voire d'une cire.

[0060] Les compositions fongicides selon l'invention peuvent être formulées sous la forme de granulés dispersables dans l'eau également compris dans le cadre de l'invention.

[0061] Ces granulés dispersables, de densité apparente généralement comprise entre environ 0,3 et 0,6 ont une dimension de particules généralement comprise entre environ 150 et 2000 et de préférence entre 300 et 1500 microns.

[0062] La teneur en matière active de ces granulés est généralement comprise entre environ 1 % et 90 %, et de préférence entre 25 % et 90 %.

[0063] Le reste du granulé est essentiellement composé d'une charge solide et éventuellement d'adjuvants tensioactifs conférant au granulé des propriétés de dispersibilité dans l'eau. Ces granulés peuvent être essentiellement de deux types distincts selon que la charge retenue est soluble ou non dans l'eau. Lorsque la charge est hydrosoluble, elle peut être minérale ou, de préférence, organique. Dans le cas d'une

charge insoluble, celle-ci est de préférence minérale, comme par exemple le kaolin ou la bentonite. Elle est alors avantageusement accompagnée d'agents tensioactifs (à raison de 0,5 à 20 % en poids du granulé) dont plus de la moitié est, par exemple, constituée par au moins un agent dispersant, essentiellement anionique, tel qu'un polynaphtalène sulfonate alcalin ou alcalino-terreux ou un lignosulfonate alcalin ou alcalino-terreux, le reste étant constitué par des mouillants non ioniques ou anioniques tel qu'un alcoylnaphtalène sulfonate alcalin ou alcalino-terreux.

[0064] Par ailleurs, bien que cela ne soit pas indispensable, on peut ajouter d'autres adjuvants tels que des agents anti-mousse.

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[0065] Le granulé selon l'invention peut être préparé par mélange des ingrédients nécessaires puis granulation selon plusieurs techniques en soi connues (drageoir, lit fluide, atomiseur, extrusion, etc.). On termine généralement par un concassage suivi d'un tamisage à la dimension de particule choisie dans les limites mentionnées ci-dessus. On peut encore utiliser des granulés obtenus comme précédemment puis imprégnés avec une composition contenant la matière active.

[0066] De manière tout à fait avantageuse, les compositions selon la présente invention sont adaptées pour une application, sur les fruits à traiter, par pulvérisation, par trempage, par pelliculage, par enrobage, etc.

[0067] La présente invention concerne également un procédé de traitement des fruits, caractérisé en ce qu'on l'on traite les fruits avec une composition fongicide selon l'invention, par trempage, pulvérisation, brossage, enrobage, pelliculage, etc. Ces techniques de traitement sont bien connues de l'homme du métier.

[0068] À titre d'exemple, le traitement par trempage consiste à tremper directement les fruits dans une solution aqueuse d'une composition fongicide selon l'invention. Cette opération peut être effectuée manuellement ou mécaniquement.

[0069] Un traitement mécanique particulièrement préféré est le brossage, c'est-à-dire que les composés fongicides sont appliqués à l'aide de brosses situées, par exemple, au-dessus de rouleaux sur lesquels se déplacent les fruits.

[0070] D'autres techniques, comme la pulvérisation au moyen d'appareils munis de buses, ou encore le pelliculage ou l'enrobage, dans des cylindres ou des "bétonnières" munis ou non de rouleaux, de brosses et/ou de pas de vis, font appel à des techniques connues qui ne seront pas développées ici.

[0071] Il est à noter que le traitement des fruits à l'aide des compositions fongicides selon l'invention est effectué de préférence après la récolte, de manière préventive et/ou curative, c'est-à-dire avant et/ou après qu'une maladie ne soit apparue sur les fruits traités.

[0072] Les composés fongicides compris dans les compositions selon l'invention peuvent être appliqués sur les fruits de manière simultanée, séquencée ou séparée.

[0073] De plus, le traitement avec les compositions fongicides selon l'invention est compatible avec un traitement pré-récolte classique, ce traitement pouvant être de tout type connu, fongicide et/ou insecticide.

[0074] Un tel traitement fongicide et/ou insecticide combiné avec un traitement, de préférence après récolte, avec une ou plusieurs compositions fongicides selon la présente invention est également compris dans le champ de la présente invention.

[0075] Enfin, la présente invention concerne également les fruits traités par une ou plusieurs compositions selon la présente invention.

[0076] Les exemples suivants sont donnés à titre non limitatifs dans le seul but de montrer l'excellente efficacité des compositions selon l'invention, et de montrer comment elles sont mises en œuvre.

Exemple 1 (action curative):

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Sur des citrons, une petite zone du zeste a été enlevée (de l'ordre de 1 cm²). Après avoir réalisé cette opération, une inoculation par une suspension de spores de *Penicillium digitatum* (concentration 3.900.000 spores/ml) à raison de 5 gouttes par fruit a été effectuée à l'endroit où le zeste a été enlevé. Les citrons, à raison de 3 fruits/facteur d'essai, ont été placés dans des coupelles, ensachés et mis en incubation à 20°C pendant 24 heures. Après ce délai, un traitement fongicide par pulvérisation a été pratiqué. Les produits et doses employées ont été les suivants :

- Composition de référence A: fénamidone + fosétyl-Al aux doses de 17 + 250, 33 + 500, 66 + 1000, 133g + 2000 mg de matières actives par litre.
- Composition de référence B: imazalil (formulation commerciale Deccosil Agrumes*, concentré émulsionnable à 200g/l) aux doses de 125, 250, 500 et 1000 mg d'imazalil par litre.

- Exemple de composition 1: Composition A + Composition B aux doses de (17+250)+125, (33+500)+250, (66+1000)+500 et (133+2000)+1000 mg de matières actives par litre (premier lot de doses).
- Exemple de composition 2: Composition A + Composition B aux doses de (10+150)+75, (20+300)+150, (40+600)+300 et (80+1200)+600 mg de matières actives par litre (second lot de doses).
- 10 [0077] Après le traitement, les citrons ont été de nouveau ensachés puis replacés à 20°C pour 6 jours.

Au 6e jour après le traitement, une notation a été effectuée. Elle a consisté à déterminer le pourcentage de surface contaminée et, par comparaison à un témoin non traité et inoculé, à définir le pourcentage d'efficacité.

Résultats:

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Composition	Doses (mg/l) (fénamidone+fosétyl-Al)+imazalil	% efficacité
Composition A	(17+250)	0
-	(33 + 500)	16
	(66+1000)	0
	(133 + 2000)	0
Composition B	125	80
_	250	32
	500	93
	1000	96
Composition 1	(17+250)+125	52
(Premier lot de doses)	(33+500)+250	88
	(66+1000)+500	93
	(133 + 2000) + 1000	98
Composition 2	(10+150)+75	60
(Second lot de doses)	(20+300)+150	88
	(40+600)+300	95
	(80 + 1200) + 600	98

Le témoin non-traité est contaminé à 83%.

Exemple 1 (action préventive):

Le protocole expérimental suivi est identique au précédent mis à part le fait que le traitement est cette fois-ci effectué un jour <u>avant</u> l'inoculation.

Résultats:

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Composition	Doses (mg/l) (fénamidone+fosétyl-Al)+imazalil	% efficacité
Composition A	(17 + 250)	0
•	(33+500)	. 11
	(66+1000)	0
	(133 + 2000)	39
Composition B	125	67
-	250	32
	500	93
	1000	100
Composition 1	(17+250)+125	87
(Premier lot de doses)	(33 + 500) + 250	92
·	(66+1000)+500	98
	(133 + 2000) + 1000	100
Composition 2	(10+150)+75	78
(Second lot de doses)	(20+300)+150	92
	(40+600)+300	94
	(80+1200)+600	100

10 Le témoin non-traité est contaminé à 30%.

[0078] Les résultats ci-dessus montrent que :

- les compositions selon l'invention possèdent une activité préventive et curative par rapport à des témoins non traités.
- les compositions selon l'invention ont une meilleure efficacité par rapport aux compositions connues, et ce même à des doses de produits comparables inférieures.

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REVENDICATIONS

- 1. Compositions fongicides utiles pour lutter contre les champignons phytopathogènes infestant ou susceptibles d'infester les fruits, caractérisée en ce qu'elles contiennent :
 - a) au moins un composé fongicide inhibiteur de la respiration mitochondriale, et
 - b) au moins un composé fongicide inhibiteur de la biosynthèse des stérols.
- 2. Compositions fongicides selon la revendication 1, caractérisées en ce que le composé fongicide inhibiteur de la respiration mitochondriale est choisi parmi l'azoxystrobine, le krésoxym-méthyl, la trifloxystrobine, la picoxystrobine, la discoxystrobine, le 4-chloro-2-cyano-N,N-dimethyl-5-p-tolylimidazole-1-sulfonamide, la famoxadone et les composés de formule générale (I):

$$(Y)_{n} \xrightarrow{CH_{1}} \overset{N}{N} \xrightarrow{M-CH_{2}} (Y)_{n}$$

$$(I)$$

dans laquelle:

- M représente un atome d'oxygène ou de soufre ;
- n est un nombre entier égal à 0 ou 1 ;
- Y est un atome de fluor ou de chlore, ou un radical méthyle.
- 3. Compositions fongicides selon la revendication 2, caractérisées en ce que le composé fongicide inhibiteur de la respiration mitochondriale est choisi parmi la famoxadone et un composé de formule (I) tel que défini dans la revendication 2.
- 4. Compositions fongicides selon l'une des revendications 2 ou 3, caractérisées en ce que le composé fongicide inhibiteur de la respiration mitochondriale est la fénamidone.
- 5. Compositions fongicides selon l'une quelconque des revendications précédentes, caractérisées en ce que le composé inhibiteur de la biosynthèse des stérols est l'imazalil.
 - 6. Compositions fongicides selon l'une quelconque des revendications

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précédentes, caractérisées en ce qu'elles comprennent l'imazalil en association avec la fénamidone.

- 7. Compositions fongicides selon l'une quelconque des revendications précédentes, caractérisées en ce que les doses de composés fongicides inhibiteurs de la respiration mitochondriale sont comprises entre 10 mg/l et 1000 mg/l, de préférence entre 20 mg/l et 300 mg/l, de préférence encore entre 40 mg/l et 150 mg/l, ou entre 50 mg/l et 100 mg/l.
- 8. Compositions fongicides selon l'une quelconque des revendications précédentes, caractérisées en ce que les doses de composés fongicides inhibiteurs de biosynthèse des stérols sont comprises entre 100 mg/l et 3000 mg/l, de préférence comprises entre 50 mg/l et 2500 mg/l, de préférence encore entre 200 mg/l et 2000 mg/l, ou entre environ 400 mg/l et 1000 mg/l.
 - 9. Compositions fongicides selon l'une quelconque des revendications précédentes, caractérisées en ce qu'elles comprennent en outre un ou plusieurs autres composés fongicides.
 - 10. Compositions fongicides selon la revendication 9, caractérisées en ce que l'autre composé fongicide est choisi parmi l'acide phosphoreux, ses dérivés et ses sels.
 - 11. Compositions fongicides selon l'une des revendications 9 ou 10, caractérisées en ce que l'autre composé fongicide est le fosétyl-Al.
 - 12. Compositions fongicides selon l'une quelconque des revendications 9 à 11, caractérisées en ce l'autre composé fongicide est présent à des doses comprises entre 500 mg/l et 6000 mg/l, par exemple entre 2000 mg/l et 4000 mg/l.
 - 13. Compositions fongicides selon l'une quelconque des revendications précédentes, caractérisées en ce qu'elles comprennent imazalil, fénamidone et fosétyl-Al.
 - 14. Compositions fongicides selon l'une quelconque des revendications précédentes, caractérisées en ce qu'elles sont utiles pour traiter un ou plusieurs des champignons phytopathogènes choisis parmi :

 Phytophthora spp., par exemple la pourriture brune des agrumes (Phytophthora parasitica), et la gommose des citrus (Phytophthora citrophthora);

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Penicillium spp., par exemple la moisissure bleue (Penicillium italicum), et la moisissure verte (Penicillium digitatum);

la pourriture amère des agrumes (Geotrichum candidum);

le black-rot des agrumes (Alternaria citri);

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l'anthracnose (Colleotrichum gloeosporioides); et

la mélanose ou pourriture phomopsienne (Diplodia natalensis ou Phomopsis citri).

- 15. Compositions fongicides selon l'une quelconque des revendications précédentes, caractérisées en ce qu'elles protègent ou luttent contre les attaques fongiques et empêchent ou enrayent le pourrissement de fruits comestibles.
- 16. Compositions fongicides selon l'une quelconque des revendications précédentes, caractérisées en ce que les fruits sont des agrumes.
- 17. Compositions fongicides selon l'une quelconque des revendications précédentes, caractérisées en ce qu'elles comprennent, outre les composés fongicides décrites dans les revendications précédentes, un ou plusieurs supports inertes solides ou liquides, agents tensioactifs, colloïdes protecteurs, adhésifs, épaississants, agents thixotropes, agents de pénétration, stabilisants, séquestrants, agents de texture, agents de saveur, exhausteurs de goût, sucres, édulcorants, et/ou des colorants.
 - 18. Compositions fongicides selon l'une quelconque des revendications précédentes, caractérisées en ce qu'elles contiennent 0,05 à 95 % en poids de matière active.
 - 19. Compositions fongicides selon l'une quelconque des revendications précédentes, caractérisées en ce qu'elles sont sous forme solide ou liquide.
 - 20. Procédé de traitement des fruits, caractérisé en ce qu'on l'on traite les fruits avec une composition fongicide selon l'une quelconque des revendications précédentes, par trempage, pulvérisation, brossage, enrobage, ou pelliculage des dits fruits.
 - 21. Procédé de traitement des fruits selon la revendication 20, caractérisé en ce que le dit traitement est effectué de manière préventive et/ou curative.
 - 22. Procédé de traitement des fruits selon la revendication 20, caractérisé en ce que le dit traitement est effectué après la récolte des fruits.

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23. Procédé de traitement des fruits selon l'une des revendications 20 à 22, caractérisé en ce que les composés fongicides sont appliqués de manière simultanée, séquencée ou séparée.

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- 24. Procédé de traitement des fruits, caractérisé en ce qu'il combine un traitement fongicide et/ou insecticide avec un traitement par une ou plusieurs compositions fongicides selon l'une quelconque des revendications 1 à 19.
- 25. Fruits traités par une ou plusieurs compositions selon la présente invention.

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A. CLASSIFICATION OF SUBJECT MATTER
IPC 7 A01N43/50 A01N57/12

A01N61/00

According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)

IPC 7 AO1N

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practical, search terms used)

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Х	EP 0 531 837 A (BASF) 17 March 1993 (1993-03-17) see the whole document	1,2,5,7, 8,14-25
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Further documents are fisted in the continuation of box C.	Patent family members are listed in annex.
"Special categories of cited documents: "A" document defining the general state of the art which is not considered to be of particular relevance "E" earlier document but published on or after the international filling date "L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified) "O" document referring to an oral disclosure, use, exhibition or other means "P" document published prior to the international filling date but later than the priority date claimed	"T" later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone "Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art. "&" document member of the same patent family
Date of the actual completion of the international search 3 April 2000	Date of mailing of the international search report $11/04/2000$
Name and mailing address of the ISA European Patent Office, P.B. 5818 Patentlaan 2 NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo nl, Fax: (+31-70) 340-3016	Authorized officer Fort, M

INTE TIONAL SEARCH REPORT

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RAPPORT DE RECHERCIT INTERNATIONALE

PCT/FR 00/00339

A. CLASSEMENT DE L'OBJET DE LA DEMANDE CIB 7 A01N43/50 A01N57/12

A01N61/00

Selon la classification internationale des brevets (CIB) ou à la fois selon la classification nationale et la CIB

B. DOMAINES SUR LESQUELS LA RECHERCHE A PORTE

Documentation minimale consultée (système de classification suivi des symboles de classement) C TB 7 A01N

Documentation consultée autre que la documentation minimale dans la mesure où ces documents relèvent des domaines sur lesquels a porté la recherche

Base de données électronique consultée au cours de la recherche internationale (nom de la base de données, et si réalisable, termes de recherche utilisés)

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Х	WO 96 03044 A (RHONE POULENC) 8 février 1996 (1996-02-08) page 1, ligne 31 -page 4, ligne 13 exemples 1,2,12,21-29	1,2,4, 7-12, 14-25
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	-/	

X Voir la suite du cadre C pour la fin de la liste des documents	Les documents de familles de brevets sont indiqués en annexe
"A" document définissant l'état général de la technique, non considére comme particulièrement pertinent "E" document antérieur, mais publié à la date de dépôt international ou après cette date "L" document pouvant jeter un doute sur une revendication de priorité ou cité pour déterminer la date de publication d'une autre citation ou pour une raison spéciale (telle qu'indiquée) "O" document se référant à une divulgation orale, à un usage, à une exposition ou tous autres moyens "P" document publié avant la date de dépôt international, mais	T" document ulterieur publié après la date de dépôt international ou la date de priorité et n'appartenenant pas à l'état de la technique pertinent, mais cité pour comprendre le principe ou la théorie constituant la base de l'invention X" document particulièrement pertinent; l'inven tion revendiquée ne peut être considérée comme nouvelle ou comme impliquant une activité inventive par rapport au document considéré isolément Y" document particulièrement pertinent; l'inven tion revendiquée ne peut être considérée comme impliquant une activité inventive lorsque le document est associé à un ou plusieurs autres documents de même nature, cette combinaison étant évidente pour une personne du métier &" document qui fait partie de la même famille de brevets
Date à laquelle la recherche internationale a été effectivement achevée 3 avril 2000	Date d'expédition du présent rapport de recherche internationale 11/04/2000
Nom et adresse postale de l'administration chargée de la recherche internationale Office Européen des Brevets, P.B. 5818 Patentlaan 2 NL – 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo nl, Fax: (+31-70) 340-3016	Font, M

RAPPORT DE RECEPCIONALE

er le Internationale No PCT/FR 00/00339

		PCI/FR 00	/ 00339
	OCUMENTS CONSIDERES COMME PERTINENTS	Ai	1
Catégorie 1	Identification des documents cités. avec le cas échéant. l'indicationdes passages per	runents	no. des revendications visees
Х	WO 93 22921 A (ZENECA) 25 novembre 1993 (1993-11-25) * voir le document entier*		1,2,7-9, 14-25
X	WO 98 33381 A (DU PONT DE NEMOURS) 6 août 1998 (1998-08-06) * voir p.12, "Preferred 6"* page 18, ligne 8 -page 19, ligne 21 page 34, ligne 6 - ligne 18 revendications 1,6		1,2,7-9, 14-25
X	"MIXTURES OF FUNGICIDES, INSECTICIDES AND HERBICIDES", RESEARCH DISCLOSURE, NR. 388, PAGE(S) 489/490 XP000635427 ISSN: 0374-4353 * voir le document entier*		1-3,7-9, 14-25
i			

RAPPORT DE RECHERCHE INTERNATIONALE

Renseignements relatifsx membrance familles de brevets

Dei ernationale No PCT/FR 00/00339

Document brevet cité au rapport de recherche	Date de publication	Membre(s) de la famille de brevet(s)	Date de publication
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Renseignements relatifs aux membres de familles de brevets

er le Internationale No PCT/FR 00/00339

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				AU	665088 B	14-12-1995
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				EP	0967869 A	05-01-2000



PATENT COOPERATION TREATY

From the INTERNATIONAL PRELIMINARY EXAMINING AUTHORITY

[stamp]

To:

BRACHOTTE Charles AVENTIS CROPSCIENCE S.A. BP 9163 69263 LYON CEDEX 09 FRANCE PCT

NOTIFICATION OF TRANSMITTAL OF INTERNATIONAL PRELIMINARY EXAMINATION REPORT

Applicant's or agent's file reference PH 99006

International application No. PCT/FR00/00339

Applicant AVENTIS CROPSCIENCE S.A. et al.

- 1. The applicant is hereby notified that this International Preliminary Examining Authority transmits herewith the international preliminary examination report and its annexes, if any, established on the international application.
- A copy of the report and its annexes, if any, is being transmitted to the International Bureau for communication to all the elected Offices.
- 3. Where required by any of the elected Offices, the International Bureau will prepare an English translation of the report (but not of any annexes) and will transmit such translation to those Offices.
- 4. REMINDER

The applicant must enter the national phase before each elected Office by performing certain acts (filing translations and paying national fees) within 30 months from the priority date (or later in some Offices) (Article 39(1)) (see also the reminder sent by the International Bureau with Form PCT/IB/301).

Where a translation of the international application must be furnished to an elected Office, that translation must contain a translation of any annexes to the International preliminary examination report. It is the applicant's responsibility to prepare and furnish such translation directly to each elected Office concerned.

For further details on the applicable time limits and requirements of the elected Offices, see Volume II of the PCT Applicant's Guide.

Name and mailing address of the IPEA/

Authorized officer:

<u>a</u>))

European Patent Office D-80298 Munich Tel. +49 89 2399-0, Tx: 523656 epmu d Fax: +49 89 2399-4465

Gallego, A

Tel. +49 89 2399-8102





PATENT COOPERATION TREATY

PCT

INTERNATIONAL PRELIMINARY EXAMINATION REPORT

(PCT Article 36 and Rule 70)

Applicant's or Agent's file reference PH 99006			s file reference	FOR FURTHER ACTION	See Notification of Transmittal of International Preliminal Examination Report (Form PCT/IPEA/416)		
	ernational a		tion No.	International filing date (11/02/2000	day/month/year)	Priority date (day/month/year) 12/02/1999	
	ernational F 1N43/50	Patent	Classification (IPC) or n	ational classification and I	PC		
	olicant ENTIS CR	OPSCI	ENCE S.A. et al.	1	· · · · · · · · · · · · · · · · · · ·		
1.			al preliminary examinate applicant according to		ared by this Internat	ional Preliminary Examining Authority and is	
2.	☐ This	repor	t is also accompanied and are the basis for th	is report and/or sheets co ive Instructions of the PCT	ts of the description	n, claims and/or drawings which have been s made before this Authority (see Rule 70.16	
	These ar	nexes	consist of a total of	sheets.			
3.	This repo	ort conf	ains indications relating	to the following items:			
	1	\boxtimes	Basis of the report				
	11		Priority				
	HI.		Non-establishment of	opinion with regard to nov	elty, inventive step a	nd industrial applicability	
	IV		Lack of unity of invent	tion			
	V	×		according to Article 35(2 tions supporting such state		elty, inventive step or industrial applicability;	
	VI		Certain documents cir	ed			
	VII	\boxtimes	Certain defects in the	international application			
	VIII	Ø	Certain observations	on the international applica	ation		
Dat	e of submi	ssion o	of the demand		Date of completion of	this report	
04/0	04/09/2000				10.05.2001		
Nar	Name and mailing address of the IPEA/ Buropean Patent Office				Authorized officer:		
	<u>9))</u>	D-80	298 Munich -49 89 2399-0, Tx: 5236	56 epmu d	Krattinger, B		
			+49 89 2399-4465		Felenhone No. ±49.8	Q 23QQ 855N	

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I.	Bas	sis of the report								
1.	by t	is report has been drawn up on the basis of the following elements (the replacement sheets received the receiving office in response to an invitation according to Article 14 are considered in the present bort as "originally filed" and are not annexed to the report as they contain no amendments (Rules 70.16 of 70.17).):								
	Des	scription, pages:								
	1-13	as originally filed								
	Clai	Claims, No.:								
	1-25	as originally filed								
2.	With in th	regard to the language , all the elements marked above were available or furnished to this Authority le language in which the international application was filed, unless otherwise indicated under this item.								
	The	se elements were available or furnished to this Authority in the following language which is:								
		the language of a translation furnished for the purposes of international search (under Rule 23.1(b)).								
		the language of publication of the international application (under Rule 48.3(b)).								
		the language of the translation furnished for the purposes of international preliminary examination (under Rule 55.2 and/or 55.3).								
3.		n regard to any nucleotide and/or amino acid sequence disclosed in the international application, international preliminary examination was carried out on the basis of the sequence listing:								
		contained in the international application in written form.								
		filed together with the international application in computer readable form.								
		furnished subsequently to this Authority in written form.								
		furnished subsequently to this Authority in computer readable form.								
		The statement that the subsequently furnished written sequence listing does not go beyond the disclosure in the international application as filed has been furnished.								
		The statement that the information recorded in computer readable form is identical to the written sequence listing has been furnished.								
4.		The amendments have resulted in the cancellation of:								
		the description, pages								
		the claims, Nos.								
		the drawings, sheets/fig								
5.		This report has been written disregarding (some of) the amendments, which were considered as going beyond the description of the invention, as filed, as is indicated below (Rule 70.2(c)):								
		(All replacement sheets comprising amendments of this nature should be indicated in point 1 and attached to this report).								

- 6. Additional observations, if necessary:
- V. Reasoned statement under Article 35(2) with regard to novelty, inventive step or industrial applicability; citations and explanations supporting such statement
- 1. Statement

Novelty

Yes:

Claims

6, 12-13

No: Claims

1-5. 7-11, 14-25

Inventive Step

Yes: No: Claims

Claims

1-25

Industrial Applicability

Yes: No: Claims Claims 1-25

2. Citations and explanations see separate sheet

VII. Certain defects in the international application

The following defects in the form or contents of the international application have been noted: see separate sheet

VIII. Certain observations in the international application

The following observations on the clarity of the claims, descriptions, and drawings or on the question whether the claims are fully supported by the description, are made: see separate sheet

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As regards point V

Reference is made to the following documents:

D1: WO 96 03044 A

D2: EP-A-0 531 837

D3: FR-A-2 754 424

D4: EP-A-0 524 496

D5: WO 93 22921 A

D6: WO 98 33381 A

D7: Research disclosure No. 388, pages

489/490.

D8: WO90/12791

Document D8 is cited in document D7. A copy of this document is attached to the present letter.

A. The present application relates to fungicide compositions which contain at least one fungicide compound inhibiting mitochondrial respiration (a) and at least one compound inhibiting sterol biosynthesis (b) (claims 1-19), methods for treating fruits with these compositions (claims 20-24) and the fruits treated with these compositions (claim 25).

Novelty

- B.1. In claim 1, the expression "useful for controlling ... fruits" is not considered as describing a technical characteristic and is not taken into account for determining novelty. Furthermore, the compositions are defined by their content, not by their use. The result is that the subject-matter of claims 14-16 is strictly the same as that of claim 1, and that these claims ought to be written as claims for using the composition and not as claims for compositions (Art. 6 PCT).
- 2. Moreover, in claims 7, 8, 12, 14, the expressions "preferably", "for example" do not have a limiting effect on the scope of the claims. Thus, the characteristic which follows such an expression is

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considered as being optional. Such expressions ought to be avoided (Art. 6 PCT).

C.1. Document D1 describes compositions comprising a compound inhibiting mitochondrial respiration of the 2-imidazolin-5-one type (= formula (I)), in particular fenamidone, and a fungicide compound inhibiting sterol biosynthesis, of the triazole, morpholine or pyrimidine type (claims 1-4, 10, examples 6, 9, 19, 21-26, 28, 29).

These compositions may contain other fungicide compounds; they comprise, in addition, solid or liquid carriers, surfactants, colloids and the like (page 8, lines 9-22), contain 0.5 to 95% by weight of active products which are in proportions of 0.001 to 2 (page 5, line 10, page 8, lines 25-26, page 9, lines 14-15); they are provided in solid or liquid form (page 9, lines 18-19).

They are applied to fruits - tomatoes, Curcubitaceae, cucumbers, cereals, wheat (page 15, line 15 to page 16, line 8) -, curatively or preventively (page 15, lines 8-9). They may be coupled with other fungicide treatments (page 2, line 9 to page 3, line 11 "at least one fungicide compound B").

These compositions are applied by spraying or daubing (page 8, line 15; page 16, lines 13-14).

Thus, in the light of the content of D1, the subject-matter of claims 1-2, 4, 7-9, 14-21, 23-25 is not novel.

The compositions of D1 do not comprise imazalil or famoxadone, and are not used for treating fruits after harvest. D1 does not specifically describe compositions comprising a compound (a), a compound (b)

and a phosphorous acid derivative (such compositions would be the result of a selection).

2. Document D2 describes the combination of a derivative inhibiting mitochondrial respiration, methyl (α-methoxyimino-2-((2-methylphenoxy)methyl))benzene acetate) (=kremoxy-methyl), with other derivatives inhibiting sterol biosynthesis, such as triazoles and imizalil (page 3, lines 1-35).

These compositions may, in addition, comprise other fungicide compounds (page 4, lines 49-51), are provided in solid or liquid form (page 4, lines 52-54), comprise solid or liquid carriers, surfactants and the like (page 5, lines 3-29), contain between 0.1 and 95% by weight of active agent in (a)/(b) ratios varying from 1:10 to 10:1 (see in particular the example).

The compositions are applied to fruits -tomatoes, apples, Curcubitaceae, cucumber, wheat, barley and the like (page 5, line 33 to page 6, line 3), by spraying (page 4, line 55), curatively or preventively (page 5, line 40) and may be combined with other insecticide or fungicide treatments (page 4, lines 49-51). They may in particular be used on seeds, that is to say on seed grains (= fruits after harvest) (page 5, line 37).

Thus, the content of document D2 affects the novelty of claims 1, 2, 5, 7-9, 14-25.

The compositions of D2 do not contain famoxadone or fenamidone or a phosphorous acid derivative.

3. Document D3 describes compositions containing a compound inhibiting mitochondrial respiration similar to strobilurin, in particular azoxystrobin, and a derivative inhibiting sterol biosynthesis, namely a

compound of the triazole type (page 1, line 27 to page 2, line 26). These compositions are provided in solid or liquid form (page 5, lines 15-16), contain solid or liquid carriers, surfactants, colloids and the like (page 4, lines 6-10, 16-21), contain 0.05 to 95% of active products by weight (page 4, lines 22-25) in ratios of between 0.01 and 30.

These compositions are used curatively or preventively (page 1, line 6), are applied by dusting, film-coating or coating (page 4, line 12; page 5, line 17, 23, page 10, line 15), are used for treating seeds (= seed grains = fruits after harvest) (page 4, lines 11; page 6, lines 33-34; page 10, lines 7-15). The application may be carried out simultaneously, sequentially or separately.

Thus in the light of the content of D3, the subject-matter of claims 1, 2, 7, 8, 14-23 and 25 is not novel.

The compositions of D3 do not contain famoxadone or fenamidone or imazalil or a phosphorous acid derivative, or additional fungicide compounds and are not coupled with other treatments.

4. Document D4 describes compositions comprising kremoxy-methyl, an inhibitor of mitochondrial respiration, with derivatives of morpholine, an inhibitor of sterol biosynthesis (page 1).

These compositions may contain other fungicide or insecticide derivatives (page 3, lines 36-38), are provided in solid or liquid form (page 3, line 40), and contain solid or liquid carriers, surfactants and the like (page 3, lines 47-56).

These compositions are sprayed (page 3, lines 39-40), applied to seeds (page 3, line 45; page 4, line

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24), fruits (page 4, lines 19-46), inter alia, Curcubitaceae, cucumbers, tomatoes, apples, cereals; they contain quantities identical to those of the present claims 7, 8 and 18 (page 2, lines 53, 54 and page 4, line 55, example).

Consequently, in the light of the content of D4, the subject-matter of claims 1, 2, 7, 8, 9, 14-25 is not novel.

The compositions of D3 do not contain famoxadone or fenamidone or imazalil or a phosphorous acid derivative.

5. Document D5 describes compositions comprising a compound inhibiting mitochondrial respiration which is similar to strobilurin, azoxystrobin and a derivative inhibiting sterol biosynthesis of the triazole type (claim 1). The compositions comprise 0.0001 to 95% of active substances by weight, in proportions ranging from 400:1 to 10:90 (page 1, lines 31-33; page 4, lines 28-30, page 4, lines 35-38), exist in solid or liquid form, and contain solid or liquid carriers, surfactants and the like (page 5, line 17 to page 7, line 29). These compositions may, in addition, comprise other fungicides or insecticides (page 8, lines 5-8). Some fungicides (page 8, line 9 to page 9, line 26) are inhibitors of sterol biosynthesis, such as triazoles the (bromuconazole. difenoconazole and like), imidazoles (imazalil), derivatives of morpholines (aldimorph, dodemorph and the like), of pyridines (buthiobate, pyrefenox), of pyrimidines (fenarimol, nuarimol and the like), of piperidines (fenpropidin). Others are of the phosphorous acid family (phosphorous tolclfos-methyl, acid, iprobenphos, pyrazophos, ampropyphos, edifenphos and phosetyl-Al (= fosetyl-Al).

These compositions are used - by spraying or coating (page 5, line 3, 25) on fruits - apples, pears,

cereals, Curcubitaceae, grapes, bananas, strawberries, tomatoes, melon, lemon and the like (page 2, line 16 to page 4, line 22) - in particular after harvesting for example for oranges (page 3, line 35), seeds (page 4, lines 25-26), curatively and preventively (page 5, lines 13-14).

Consequently, in the light of the content of D5, the subject-matter of claims 1, 2, 5, 7-11, 14-25 is not novel.

The compositions of D5 do not contain famoxadone or fenamidone. D5 does not specify the quantity of additional fungicide.

Document D6 describes compositions based on quinazolinone and derivatives inhibiting biosynthesis, including imazalil (see list pages compounds inhibiting 18, 19) mitochondrial respiration including azoxystrobin, kremoxym-methyl and another strobilurin derivative. They are provided in liquid or solid form, contain liquid or solid carriers, surfactants and the like (page 4, lines 13-22; page 18, lines 3-7; pages 30-31). They may contain other insecticides and fungicides (page 33), including phosphorous acid derivatives, such as edifenphos, iprobenphos and fosetyl-aluminium (page 33, line 26 to 35). The percentage of active ingredient varies from 0.01 to 99% (page 30), and the proportion of compounds (a)/(b) varies from 1:30 to 30:1 (page 4, line 22-27).

These compositions are applied to fruits, seeds (page 4, line 12; page 32, lines 29, 33), curatively and preventively (page 34, lines 7-11).

If the quinazolinones of D6 have an inhibitory action on mitochondrial respiration or sterol biosynthesis, then in the light of the content of D6, the subject-matter of claims 1, 2, 5, 7-11, 14-25 is

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not novel. The Applicant is invited to specify the mode of action of the quinazolinones.

Document D6 does not disclose a composition based on famoxadone, on fenamidone and does not specify the quantity of additional fungicide agent.

7. Document D7 describes compositions comprising famoxadone and other fungicide compounds inhibiting sterol biosynthesis, such as triazoles (bromuconazole, cyproconazole, imidazoles (imazalil, prochloraz and the like), pyrimidines (fenarimol, nuarimol and the like), pyridines (pyrefenox), morpholines (dodemorph, fenpropimorph and the like), piperidines (fenpropidin), thus affecting the novelty of the subject-matter of claims 1-3. These compositions are provided in solid or liquid form and optionally contain surfactants (lines 1-10).

Thus, in the light of the content of D7, the subject-matter of claims 1-3, 5, 14-17, 19 is not novel.

Document D7 does not describe the proportions and the quantities of active products, does not specify the methods of application or the subjects targeted by the application. Furthermore, it does not describe a composition specifically comprising in addition to compounds (a) and (b) a third compound (such compositions would be the result of a selection). In addition, the document does not describe compositions comprising fenamidone.

8. Document D8 describes compositions comprising famoxadone and other fungicide compounds inhibiting sterol biosynthesis (pages 8-9, 46), such as triazoles (bromuconazole, tebuconazole and the like), morpholines (fenpropimorph, page 81, line 17-19 and the like), piperidines (fenpropidin), thus affecting the novelty

of the subject-matter of claims 1-3 (page 79-81). These compositions are provided in solid or liquid form and optionally contain surfactants (page 75), contain 1 to 99% of active substance. They are used on the seeds and the fruits curatively or preventively (page 79, page 83, lines 32-38), and are applied by spraying (page 75).

Thus, in the light of the content of D8, the subject-matter of claims 1-3, 14-21, 23-25 is not novel.

Document D8 does not describe a composition specifically comprising, in addition to compounds (a) and (b), a third compound (such compositions would be the result of a selection). In addition, the document does not describe compositions comprising fenamidone or imazalil.

9. In claim 25, the fruits treated with the compositions of the inventions are not different from the point of view of their structures from the untreated fruits or from the fruits treated with other plant-protection compositions, in particular fungicides. Consequently, the subject-matter of claim 25 is not novel (Art. 33(2) PCT).

10. Consequence

In the light of the preceding text, the subject-matter of claims 1-5, 7-11, 14-25 is not novel, while the subject-matter of claims 6, 12 and 13 is novel (Art. 33(2) PCT).

D. Inventive step

Documents D1-D8 which are all considered as the closest prior art all describe compositions comprising (a) at least one "fungicide compound inhibiting

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mitochondrial respiration and at least one compound inhibiting sterol biosynthesis (see objections novelty). Furthermore, these documents demonstrate the synergistic effect obtained or the efficacy of these compositions in the case of controlling fungi, particular fungi which affect fruits (D1: page 4, lines 17-25, examples; D2: page 4, lines 21-23, example; D3: page 1, lines 12-28; examples; D4: examples; D5: page 1, line 15, example; D6: examples; D7: line 13).

Consequently, the subject-matter now claimed being identical to the disclosures of the prior art documents, the advantages of the compositions being already known, the subject-matter of the present application does not involve an inventive step (Art. 33(3)PCT).

The subject-matter of the claims ought to be restricted to preferred forms of the invention which are novel and inventive in relation to the disclosures of the prior art. Demonstration by submissions or evidence that the subject-matter of the application involves an inventive step in relation to the prior art compositions would be necessary.

As regards point VII

Contrary to what is required by rule 5.1 a) ii) the description does not indicate the relevant prior state of the art disclosed in documents D1-D7 and does not cite these documents.

As regards point VIII

A discrepancy appears between the subjectmatter of claim 2 and its basis in the description: the description defines as compound а inhibiting mitochondrial respiration "strobilurin and analogous compounds or derivatives" whereas the claim only

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defines a few specific analogues, thus making the subject-matter of the claim obscure (page 3, line 9) (Art. 6 PCT).

Furthermore, the expression "strobilurin and analogous compounds or derivatives" is vague since it does not make it possible to precisely define, in terms of structures, the compounds included under this term, thus making the description vague and therefore the subject-matter of the claim obscure (Art. 6 PCT).

An additional discrepancy appears between the subject-matter of claim 2 and its basis in the description: the description defines as a compound inhibiting mitochondrial respiration "discostrobin" (page 3, line 11) whereas the claim defines "discoxystrobin", thus making the subject-matter of the claim obscure (page 3, line 9). IPEA has not found the chemical formula for this compound. The Applicant is invited to specify it.

- 2. A discrepancy appears between the subject-matter of claim 9 and its basis in the description on page 4, paragraph 33. The description defines fungicides suitable for treatments of fungal diseases of fruits, whereas the claim does not contain this characteristic, thus making the subject-matter of the claim obscure (Art. 6 PCT).
- 3. The subject-matter of claim 23 is inconsistent with the subject-matter of claims 1-19 and 20. The subject-matter of claims 1-19 consists of a composition comprising one or more compounds (a), one or more compounds (b) and optionally a third type of fungicide. If these various compounds are applied sequentially or separately, they cannot form part of the same composition and this manner of applying the products does not therefore form part of the invention as defined in claims 1-22, 24 and 25, thus making the

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scope of the set of claims obscure (Art. 6 PCT). The products (a), (b) and the optional other fungicides should form part of one and the same composition and can only be applied simultaneously in the context of the present invention.

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RAPPORT D'EXAMEN PRELIMINAIRE INTERNATIONAL

(article 36 et règle 70 du PCT)

Référence du dossier du déposant ou du mandataire PH 99006		POUR SUITE A DO			cation de transmission du rapport d'examen international (formulaire PCT/IPEA/416)		
Demande in	ntemationale n°	Date du dépot international (jour/mois/année) Date de priorité (jour/mois/année)		Date de priorité (jour/mois/année)			
PCT/FRC	00/00339	11/02/2000			12/02/1999		
	Classification internationale des brevets (CIB) ou à la fois classification nationale et CIB A01N43/50						
Déposant							
AVENTIS	CROPSCIENCE S.A. et al	l.					
	 Le présent rapport d'examen préliminaire international, établi par l'administaration chargée de l'examen préliminaire international, est transmis au déposant conformément à l'article 36. 						
2. Ce R	APPORT comprend 11 feuilles	s, y compris la présente	feuille de d	couverture			
éi l'a ad	 Il est accompagné d'ANNEXES, c'est-à-dire de feuilles de la description, des revendications ou des dessins qui ont été modifiées et qui servent de base au présent rapport ou de feuilles contenant des rectifications faites auprès de l'administration chargée de l'examen préliminaire international (voir la règle 70.16 et l'instruction 607 des Instructions administratives du PCT). Ces annexes comprennent feuilles. 						
3. Le pré	3. Le présent rapport contient des indications relatives aux points suivants:						
1	Base du rapport						
	Priorité	W. 22.2					
1111		gopinion quant a ia no	ouveaute, r	activite inv	entive et la possibilite		
IV	☐ Absence d'unité de l'inv	ention					
V	 Déclaration motivée sele d'application industrielle 				ité inventive et la possibilité léclaration		
VI	☐ Certains documents cité	ės					
VII	Irrégularités dans la der						
VIII	○ Observations relatives à	a la demande internation	nale				
Date de présentation de la demande d'examen préliminaire internationale		n préliminaire	Date d'ach	èvement du	présent rapport		
04/09/2000			10.05.2001	I			
	esse postale de l'administration cha éliminaire international:	argée de	Fonctionna	ire autorisé	JASONES MITEINE		
Office européen des brevets D-80298 Munich Tél. +49 89 2399 - 0 Tx: 523656 epmu d Fax: +49 89 2399 - 4465			Krattinge		A COO O O S S O O O O O O O O O O O O O O		
			N° de télép	hone +49 89	9 2399 8550		





Demande internationale n° PCT/FR00/00339

I. Bas du rapport

1.	à l'e rap	n ce qui concerne les éléments de la demande internationale (<i>les feuilles de remplacement qui ont été remises</i> l'office récepteur en réponse à une invitation faite conformément à l'article 14 sont considérées dans le présen apport comme "initialement déposées" et ne sont pas jointes en annexe au rapport puisqu'elles ne contiennent as de modifications (règles 70.16 et 70.17)):						
	Des	scription, pages:						
	1-1	3	version initiale					
	Rev	vendications, N°:						
	1-2	5 v	version initiale					
2.	lui c		ngue , tous les éléments indiqués ci-dessus étaient à la disposition de l'administration ou langue dans laquelle la demande internationale a été déposée, sauf indication contraire					
	Ces	Ces éléments étaient à la disposition de l'administration ou lui ont été remis dans la langue suivante: , qui est :						
		la langue d'une trad	uction remise aux fins de la recherche internationale (selon la règle 23.1(b)).					
		la langue de publica	ition de la demande internationale (selon la règle 48.3(b)).					
		la langue de la tradu 55.3).	uction remise aux fins de l'examen préliminaire internationale (selon la règle 55.2 ou					
3.	inte		séquences de nucléotides ou d'acide aminés divulguées dans la demande néant), l'examen préliminaire internationale a été effectué sur la base du listage des					
		contenu dans la der	nande internationale, sous forme écrite.					
		déposé avec la demande internationale, sous forme déchiffrable par ordinateur.						
		remis ultérieurement à l'administration, sous forme écrite.						
		remis ultérieurement à l'administration, sous forme déchiffrable par ordinateur.						
			n laquelle le listage des séquences par écrit et fourni ultérieurement ne va pas au-delà le dans la demande telle que déposée, a été fournie.					
			n laquelle les informations enregistrées sous déchiffrable par ordinateur sont identiques à es séquences Présenté par écrit, a été fournie.					
4.	Les	modifications ont en	traîné l'annulation :					
		de la description,	pages:					
		des revendications,	n ^{os} :					
		des dessins,	feuilles:					



RAPPORT D'EXAMEN PRÉLIMINAIRE INTERNATIONAL

Demande internationale n° PCT/FR00/00339

5.	Le présent rapport a été formulé abstraction faite (de certaines) des modifications, qui ont été considérées
	comme allant au-delà de l'exposé de l'invention tel qu'il a été déposé, comme il est indiqué ci-après (règle
	70.2(c)):

(Toute feuille de remplacement comportant des modifications de cette nature doit être indiquée au point 1 et annexée au présent rapport)

- 6. Observations complémentaires, le cas échéant :
- V. Déclaration motivée selon l'article 35(2) quant à la nouveauté, l'activité inventive et la possibilité d'application industrielle; citations et explications à l'appui de cette déclaration
- 1. Déclaration

Nouveauté Oui : Revendications 6, 12-13

Non: Revendications 1-5.7-11, 14-25

Activité inventive Oui : Revendications

Non: Revendications 1-25

Possibilité d'application industrielle Oui : Revendications 1-25

Non: Revendications

2. Citations et explications voir feuille séparée

VII. Irrégularités dans la demande internationale

Les irrégularités suivantes, concernant la forme ou le contenu de la demande internationale, ont été constatées : voir feuille séparée

VIII. Observations relatives à la demande internationale

Les observations suivantes sont faites au sujet de la clarté des revendications, de la description et des dessins et de la question de savoir si les revendications se fondent entièrement sur la description : voir feuille séparée

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Conc rnant le point V

Il est fait référence aux documents suivants:

D1: WO 96 03044 A

D2: EP-A-0 531 837

D3: FR-A-2 754 424

D4: EP-A-0 524 496

D5: WO 93 22921 A

D6: WO 98 33381 A

D7: Research disclosure, nr. 388, pages 489/490.

D8: WO90/12791

Le document D8 est cité dans le document D7. Une copie de ce document est jointe à la présente lettre.

La présente demande concerne des compositions fongicides qui contiennent au moins un composé fongicide inhibiteur de la respiration mitochondriale (a) et au moins un composé inhibiteur de la biosynthèse de stérols (b) (revendications 1-19), des procédés de traitement des fruits avec ces compositions (revendications 20-24) et les fruits traités par ces compositions (revendications 25).

Nouveauté

- B.1. Dans la revendications 1, l'expression "utiles pour lutter...fruits" n'est pas considérée comme décrivant une caractéristique technique et n'est pas prise en compte pour juger de la nouveauté. De plus, les compositions sont définies par leur contenu, non par leur utilisation. Il en résulte que l'objet des revendications 14-16 est strictement le même que celui de la revendication 1, et que ces revendications devraient être rédiger comme des revendications d'utilisation de la composition et non comme des revendications de compositions (Art. 6 PCT).
- 2. Par ailleurs, dans les revendications 7, 8, 12, 14, les expressions "de préférence", "par exemple" n'ont pas d'effet limitatif sur la portée des revendications. Ainsi la caractéristique qui suit une telle expression est considérée comme facultative. De telles expressions devraient être évitées (Art. 6 PCT).

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C.1. Le document D1 décrit des compositions comprenant un composé inhibiteur de la respiration mitochondriale de type 2-imidazoline-5-one (= formule (I)), en particulier la fénamidone, et un composé fongicide inhibiteur de la biosynthèse des stérols, du type triazole, morpholine, pyrimidine (revendications 1-4, 10, exemples 6, 9, 19, 21-26, 28, 29).

Ces compositions peuvent contenir d'autres composés fongicides; elles comprennent en outre des supports solides ou liquides, des tensioactifs, des colloïdes, etc. (page 8, lignes 9-22), contiennent 0.5 à 95% en poids de produits actifs qui sont dans des proportions 0.001 à 2 (page 5, lignes 10; page 8, lignes 25-26, page 9, lignes 14-15); elles se présentent sous forme solide ou liquide (page 9, lignes 18-19).

Elles sont appliquées sur des fruits - les tomates, curcubitacées, concombres, céréales, blé (page 15, ligne 15 à page 16, lignes 8)-, à titre curatif ou préventif (page 15, lignes 8-9). Elles peuvent être couplées à d'autres traitements fongicides (page 2, ligne 9 à page 3, ligne 11 "au moins un composé fongicide B")

Ces compositions sont appliquées par pulvérisation ou badigeonnage (page 8, ligne 15; page 16, lignes 13-14).

Ainsi, au vu du contenu de D1, l'objet des revendications 1-2, 4, 7-9, 14-21, 23-25 n'est pas nouveau.

Les compositions de D1 ne comprennent pas d'imazalil ou de famoxadone, et ne sont pas utilisées pour traiter les fruits après récolte. D1 ne décrit pas spécifiquement de compositions comprenant un composé (a), un composé (b) et un dérivé d'acide phosphoreux (de telles compositions seraient le résultat d'une sélection).

2. Le document D2 décrit l'association d'un dérivé inhibiteur de la respiration mitochondriale, le methyl (α-methoxyimino-2-((2-methylphenoxy)methly))benzene acetate) (= krémoxy-méthyl), avec d'autres dérivés inhibiteurs de la biosynthèse des stérols, tels les triazoles et l'imizalil (page 3, lignes 1-35).

Ces compositions peuvent en outre comprendre d'autres composés fongicides (page 4, lignes 49-51), se présentent sous forme solide ou liquide (page 4, lignes 52-54), comprennent des supports solides ou liquides, des agents tensioactifs, etc. (page 5, lignes 3-29), contiennent entre 0,1 et 95% en poids d'agent actif dans des proportions (a)/(b) variant de 1:10 à 10:1 (voir notamment l'exemple).

Les compositions sont appliquées sur des fruits -tomates, pommes,

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curcubitacées, concombre, blé, orge, etc. ((page 5, ligne 33 à page 6, ligne 3), par pulvérisation (page 4, ligne 55), à titre curatif ou préventif (page 5, ligne 40) et peuvent être combinées avec d'autres traitements insecticides ou fongicides (page 4, lignes 49-51). Elles peuvent être notamment utilisées sur les semences, c'est à dire sur les graines (= fruits après récolte) (page 5, lignes 37)

Ainsi le contenu du document D2 affecte la nouveauté des revendications 1, 2, 5, 7-9, 14-25.

Les compositions de D2 ne contiennent pas de famoxadone, ni de fénamidone, ni de dérivé de l'acide phosphoreux.

3. Le document D3 décrit des compositions contenant un composé inhibiteur de la respiration mitochondriale analogue de la strobilurine, en particulier l'azoxystrobine, et un dérivé inhibiteur de la biosynthèse des stérols, à savoir un composé de type triazole (page 1, lignes 27 à page 2, ligne 26). Ces compositions se présentent sous forme solide ou liquide (page 5, lignes 15-16), contiennent des supports solides ou liquides, des tensio-actifs, des colloïdes... (page 4, lignes 6-10, 16-21), contiennent 0.05 à 95% de produits actifs en poids (page 4, lignes 22-25) dans des rapport compris entre 0.01 et 30.

Ces compositions sont utilisées à titre curatif ou préventif (page 1, ligne 6), sont appliquées par poudrage, pelliculage ou enrobage (page 4, ligne 12; page 5, ligne 17, 23, page 10, ligne 15), sont utilisées pour traiter les semences (= graines = fruits après récoltes) (page 4, lignes 11; page 6, lignes 33-34; page 10, lignes 7-15). L'application peut être simultanée, séquentielle ou séparée.

Ainsi au vu du contenu de D3, l'objet des revendications 1, 2, 7, 8, 14-23 et 25 n'est pas nouveau.

Les compositions de D3 ne contiennent ni famoxadone, ni fénamidone, ni imazalil, ni dérivé d'acide phosphoreux, ni de composés fongicides supplémentaires et ne sont pas couplées à d'autres traitements.

4. Le document D4 décrit des compositions comprenant le krémoxy-méthyl, inhibiteur de la respiration mitochondriale, avec des dérivés de la morpholine, inhibiteur de la biosynthèse des stérols (page 1).

Ces compositions peuvent contenir d'autres dérivés fongicides ou insecticides

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(page 3, lignes 36-38), se présentent sous forme solide, liquide (page 3, ligne 40), contiennent des supports solides, liquides, des tensio actifs, ... (page 3, lignes 47-56),

Ces compositions sont pulvérisées (page 3, lignes 39-40), appliquées à des semences (page 3, ligne 45; page 4, ligne 24), des fruits (page 4, lignes 19-46), entre autre les curcubitacées, les concombres, les tomates, les pommes, les céréales; elles contiennent des quantités identiques à celles des présentes revendications 7, 8 et 18 (page 2, lignes 53, 54 et page 4, ligne 55, exemple).

Par conséquent, au vu du contenu de D4, l'objet des revendications 1, 2, 7, 8, 9, 14-25 n'est pas nouveau.

Les compositions de D3 ne contiennent ni famoxadone, ni fénamidone, ni imazalil, ni dérivé d'acide phosphoreux.

5. Le document D5 décrit des compositions comprenant un composé inhibiteur de la respiration mitochondriale analogue de la strobilurine, l'azoxystrobine et un dérivé inhibiteur de la biosynthèse des stérols de type triazole (revendication 1). Les compositions comprennent 0.0001 à 95% de matières actives en poids, dans des proportions variant de 400:1 à 10:90 (page 1, lignes 31-33; page 4, lignes 28-30, page 4, lignes 35-38), se présentent sous forme solide, liquide, contiennent des supports solides, liquides, des tensioactifs... (page 5, ligne 17 à page 7, ligne 29). Ces compositions peuvent en outre comporter d'autres fongicides ou insecticides (page 8, lignes 5-8). Certains des fongicides (page 8, lignes 9 à page 9, lignes 26), sont des inhibiteurs de la biosynthèses des stérols, comme les triazoles (bromuconazole, difenoconazole...), les imidazoles (imazalil), les dérivés de morpholines (aldimorph, dodemorph...), de pyridines (buthiobate, pyrefenox), de pyrimidines (fenarimol, nuarimol..), de pipéridines (fenpropidin). D'autres sont de la famille de l'acide phosphoreux (phosphorus acid, iprobenphos, pyrazophos, tolclfos-methyl, ampropypphos, edifenphos et phoselyl-Al (= fosetyl-Al).

Ces compositions sont utilisées - par pulvérisation ou enrobage (page 5, ligne 3, 25) sur des fruits - pommes, poires, céréales, curcubitacées, raisins, bananes, fraises, tomates, melon, citron etc. (page 2, ligne 16 à page 4, ligne 22)- notamment après récolte par exemple pour les oranges, (page 3, lignes 35), les graines (page 4, lignes 25-26), à titre curatif et préventif (page 5, lignes 13-14).

Par conséquent, au vu du contenu de D5, l'objet des revendications 1, 2, 5, 7-11,

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14-25 n'est pas nouveau.

Les compositions de D5 ne contiennent ni famoxadone, ni fénamidone. D5 ne précise pas la quantité de fongicide supplémentaire.

Le document D6 décrit des compositions à base de guinazolinone et de dérivés 6. inhibiteur de la biosynthèse des stérols, dont l'imazalil (voir liste pages 18, 19) ou de composés inhibiteurs de la respiration mitochondriale dont l'azoxystrobine, le krémoxym-méthyl et un autre dérivé de la strobilurine. Elles se présentent sous forme liquide ou solide, contiennent des supports liquides, solides, des tensioactifs, etc. (page 4, lignes 13-22; page 18, lignes 3-7; pages 30-31). Ils peuvent contenir d'autres insecticides et fongicides (page 33), dont des dérivés d'acide phosphoreux, comme edifenphos, iprobenphos et le fosétyl-aluminium (page 33, ligne 26 à 35). Le pourcentage d'ingrédient actif varie de 0.01 à 99% (page 30), et la proportion des composés (a)/(b) varie de 1:30 à 30:1 (page 4, ligne 22-27).

Ces compositions sont appliquées sur les fruits, les graines (page 4, ligne 12; page 32, lignes 29, 33), à titre curatif et préventif (page 34, lignes 7-11)

Si les quinazolinone de D6 ont une action inhibitrice de la respiration mitochondriale ou de la biosynthèse des stérols, alors au vu du contenu de D6, l'objet des revendications 1, 2, 5, 7-11, 14-25 n'est pas nouveau. Le Demandeur est invité à préciser le mode d'action des quinazolinones.

Le document D6 ne divulgue pas de composition à base de famoxadone, de fénamidone et ne précise pas la quantité de l'agent fongicide supplémentaire.

7. Le document D7 décrit des compositions comprenant la famoxadone et d'autres composés fongicides inhibiteurs de la biosynthèse des stérols, tels les triazoles (bromuconazole, cyproconazole, les imidazoles (imazalil, prochloraz..), les pyrimidines (fenarimol, nuarimol..), les pyridines (pyrefenox), les morpholines (dodemorph, fenpropimorph...), les pipéridines (fenpropidin), affectant ainsi la nouveauté de l'objet des revendications 1-3. Ces compositions se présentent sous forme solide, liquide et contiennent éventuellement des tensio-actif (lignes 1-10)

Ainsi, au vu du contenu de D7, l'objet des revendications 1-3, 5, 14-17, 19 n'est pas nouveau.

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Le document D7 ne décrit pas les proportions et les quantités de produits actifs, ne précisent pas les procédés d'application, ni les objets visés par l'application. De plus il ne décrit pas de composition comprenant spécifiquement en plus des composés (a) et (b) un troisième composé (de telles compositions seraient le résultat d'une sélection). En outre le document ne décrit pas de compositions comprenant de fénamidone.

8. Le document D8 décrit des compositions comprenant la famoxadone et d'autres composés fongicides inhibiteurs de la biosynthèse des stérols (pages 8-9, 46), tels les triazoles (bromuconazole, tebuconazole...), les morpholines (fenpropimorph, page 81, ligne 17-19...), les pipéridines (fenpropidin), affectant ainsi la nouveauté de l'objet des revendications 1-3 (page 79-81). Ces compositions se présentent sous forme solide, liquide et contiennent éventuellement des tensio-actifs (page 75), contiennent 1 à 99% de matière active. Elles sont utilisées sur les graines et les fruits à titre curatif ou préventif (page 79, page 83, lignes 32-38), et sont appliquées par pulvérisation (page 75).

Ainsi, au vu du contenu de D8, l'objet des revendications 1-3, 14-21, 23-25 n'est pas nouveau.

Le document D8 ne décrit pas de composition comprenant spécifiquement en plus des composés (a) et (b) un troisième composé (de telles compositions seraient le résultat d'une sélection). En outre le document ne décrit pas de compositions comprenant de fénamidone ou l'imazalil.

9. A la revendication 25, les fruits traités par les compositions de l'inventions ne différent pas du point de vue de leur structures des fruits non traités ou des fruits traités avec d'autres compositions phytosanitaires, en particulier, fongicides. Par conséquent l'objet de la revendication 25 n'est pas nouveau (Art. 33(2) PCT).

10. Bilan

Au vu de ce qui précède l'objet des revendications 1-5, 7-11, 14-25 n'est pas nouveau, alors que l'objet des revendications 6, 12 et 13 est nouveau (Art. 33(2) PCT,

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D. Activit inv ntiv

Les documents D1-D8 qui sont tous considérés comme l'art antérieur le plus proche, décrivent tous des compositions comprenant (a) au moins un composé fongicide inhibiteur de la respiration mitochondriale et au moins un composé inhibiteur de la biosynthèse de stérols (voir objections de nouveauté). De plus ces documents démontrent l'effet synergique obtenu ou l'efficacité de ces compositions lorsqu'il s'agit de lutter contre les champignons, en particulier les champignons affectant les fruits (D1: page 4, lignes 17-25, exemples; D2: page 4, lignes 21-23, exemple; D3: page 1, lignes 12-28; exemples; D4: exemples; D5: page 1, ligne 15, exemple; D6: exemples; D7: ligne 13).

Par conséquent, l'objet présentement revendiqué étant identique aux divulgations des documents de l'art antérieur, les avantages des compositions étant déjà connues, l'objet de la présente demande n'implique pas d'activité inventive (Art. 33(3) PCT).

L'objet des revendications devraient être restreint à des formes préférées de l'invention qui soient nouvelles et inventives par rapport aux révélations de l'art antérieur. La demonstration par argumentation ou évidence que l'objet de la future demande implique une activité inventive par rapport aux compositions de l'art antérieur serait nécessaire.

Concernant le point VII

Contrairement à ce qu'exige la règle 5.1 a) ii) PCT, la description n'indique pas l'état de la technique antérieure pertinent exposé dans les documents D1-D7 et ne cite pas ces documents.

Concernant le point VIII

Une disconcordance apparaît entre l'objet de la revendication 2 et son fondement dans la description : la description définit comme composé inhibiteur de la respiration mitochondriale "la strobilurine et dérivés ou composés analogues" alors que la revendications ne définit que quelques analogues précis, rendant ainsi l'objet de la revendication obscure (page 3, ligne 9) (Art. 6 PCT).

De plus l'expression "la strobilurine et dérivés ou composés analogues" est vague

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puisqu'elle ne permet pas de définir précisément en terme de structures les composés compris sous ce terme, rendant ainsi la description vague et donc l'objet de la revendication obscure (Art. 6 PCT).

Une disconcordance supplémentaire apparaît entre l'objet de la revendication 2 et son fondement dans la description : la description définit comme composé inhibiteur de la respiration mitochondriale "la discostrobine" (page 3, ligne 11) alors que la revendication définit la "discoxystrobine", rendant ainsi l'objet de la revendication obscure (page 3, ligne 9). L'IPEA n'as pas trouvée quelle était la formule chimique de ce composé. Le Demandeur est invité à la préciser.

- 2. Une disconcordance apparaît entre l'objet de la revendication 9 et son fondement dans la description à la page 4, paragraphe 33. La description définit des fongicides adaptés aux traitements des maladies fongicides du fruits, alors que la revendication ne contient pas cette caractéristique, rendant ainsi l'objet de la revendication obscure (Art. 6 PCT).
- L'objet de la revendication 23 est incohérent avec l'objet des revendications 1-19 et 20. L'objet des revendications 1-19 consiste en une composition comprenant un ou des composés (a), un ou des composés (b) et éventuellement un troisième type de fongicide. Si ces différents composés sont appliqués de manière séquencé ou séparée, ils ne peuvent faire partie de la même composition et cette manière d'appliquer les produits ne fait donc pas partie de l'invention telle que définie aux revendications 1-22, 24 et 25, rendant ainsi la portée du jeu de revendication obscure (Art. 6 PCT). Les produits (a), (b) et les éventuels autres fongicides doivent faire partie d'une seule et même composition et ne peuvent être appliqués que simultanément dans le cadre de la présente invention.

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RAPPORT DE RECHERCHE INTERNATIONALE

(article 18 et règles 43 et 44 du PCT)

Référence du dossier du déposant ou du mandataire	formul		mission du rapport d et, le cas échéant, le	le recherche International point 5 cl-après
PH 99006	A DONNER	·		
Demande internationale n°	Date du dépôt internations	l(jour/mois/année)	(Date de priorité (l. (jour/mois/année)	a plus ancienne)
PCT/FR 00/00339	11/02/20	00	12	/02/1999
Déposant				
AVENTIC OPODOSTENOE C A	.1 .9			
AVENTIS CROPSCIENCE S.A.	et al.			
Le présent rapport de recherche internation				ale, est transmis au
déposant conformément à l'article 18. Une	opie en est transmise au	Jureau Internationa	L.	1
Ce rapport de recherche internationale co	mprend3	feuilles.		
Il est aussi accompagné d	l'une copie de chaque docui	nent relatif à l'état c	le la technique qui y	est cité.
1. Base du rapport				
a. En ce qui concerne la langue , la langue dans laquelle elle a été dé				internationale dans la
la recherche International	e a été effectuée sur la base	d'une traduction de	e la demande interna	ationale remise à l'administration.
b. En ce qui concerne les séquence la recherche internationale a été e		ge des séquences :		de Internationale (le cas échéant),
I 😕	e internationale, sous forme		inateur.	•
	dministration, sous forme éc	•		
remis ultérleurement à l'a	dministration, sous forme dé	chiffrable par ordina	ateur.	
La déclaration, selon laqu divulgation faite dans la d	elle le listage des séquence emande telle que déposée, :	s présenté par écrit a été foumle.	et fourni uitérieuren	nent ne vas pas au-delà de la
La déclaration, selon laqu du listage des séquences	elle les informations enregis présenté par écrit, a été fou	trées sous forme dé mie.	échiffrable par ordina	ateur sont identiques à celles
2. 📗 II a été estimé que certa	ines revendications ne po	ıvaient pas faire l'	objet d'une recher	che (voir le cadre I).
3. Il y a absence d'unité de	l'invention (voir le cadre II	١.		
4. En ce qui concerne le titre,				
	u'il a été remis par le dépos			
Le texte a été établi par l'a	administration et a la teneur	sulvante:		
5. En ce qui concerne l'abrégé,				
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RAPPORT DE RECHERCHE INTERNATIONALE

PC R 00/00339

A. CLASSEMENT DE L'OBJET DE LA DEMANDE CIB 7 A01N43/50 A01N57/12 A01N61/00

Selon la classification internationale des brevets (CIB) ou à la fois selon la classification nationale et la CIB

B. DOMAINES SUR LESQUELS LA RECHERCHE A PORTE

Documentation minimale consultée (système de classification suivi des symboles de classement) CIB 7 A01N

Documentation consultée autre que la documentation minimale dans la mesure où ces documents relèvent des domaines sur lesquels a porté la recherche

Base de données électronique consultée au cours de la recherche internationale (nom de la base de données, et si réalisable, termes de recherche utilisée)

C. DOCUMENTS CONSIDERES COMME PERTINENTS				
Catégorie °	Identification des documents cités, avec, le cas échéant, l'indication des passages pertinents	no, des revendications visées		
X	W0 96 03044 A (RHONE POULENC) 8 février 1996 (1996-02-08) page 1, ligne 31 -page 4, ligne 13 exemples 1,2,12,21-29	1,2,4, 7-12, 14-25		
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Yolr la suite du cadre C pour la fin de la liste des documents	Les documents de familles de brevets sont indiqués en annexe
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Interpolication Application No
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Mixtures of Fungicides, Insecticides and Herbicides

The compound 5-methyl-5-(4-phenoxyphenyl)-3-phenylamino-2,4-oxazolidinedione which has Formula I:

is described in WO 90/12791 together with its preparation, use as a fungicide and fungicidal compositions containing it. Other methods for the preparation of the compound of Formula I are described in WO 94/11359.

The compound of Formula I may be used as a fungicide in its own right or it may be used as a fungicide in the form of a composition comprising a solid or liquid carrier or diluent and optionally a wetting, dispersing or emulsifying agent. It may also be used with one or more other fungicides, with one or more insecticides, and/or with one or more herbicides.

The compound of Formula I may be used in any proportion with other fungicides. By including another fungicide, the resulting composition can have a broader spectrum of activity, a greater level of intrinsic activity than the compound alone, or a greater level of activity than the additive control of the individual components of the composition. Examples of fungicidal compounds which may be included are: acypetacs, anilazine, azaconazole, azoxystrobin (ICIA5504), benalaxyl, benomyl, biphenyl, bitertanol, blasticidin-S, borax, Bordeaux mixture (tribasic copper sulfate), bromuconazole, bupirimate, butylamine, calcium polysulfide, captafol, captan, carbendazim, carboxin, chinomethionat, chloroneb, chlorothalonil, chlozolinate, copper hydroxide, copper oxychloride, copper salts, copper sulfate, cuprous oxide, cymoxanil, cyproconazole, cyprodinil (CGA 219417), dazomet, debacarb, dichlofluanid, dichlone, dichlorophen, diclomezine, dicloran, diethofencarb, difenoconazole, dimethirimol, dimethomorph, diniconazole, diniconazole-M, dinocap, diphenylamine, dithianon, DNOC, dodemorph, dodine, edifenphos, epoxiconazole (BAS 480F), ethirimol, ethoxyquin, etridiazole, fenarimol, fenbuconazole, fenfuram, fenpiclonil, fenpropidin, fenpropimorph, fentin, ferbam, ferimzone, fluazinam, fludioxonil, fluoromide, fluquinconazole, flusilazole, flusulfamide, flutolanil, flutriafol, folpet, formaldehyde, fosetyl-aluminum, fuberidazole, furalaxyl, guazatine, GY-81, hexachlorobenzene, hexaconazole, 8-hydroxyquinoline sulfate, hymexazol, ICIA0858, imazalil, imibenconazole, iminoctadine, ipconazole, iprobenfos, iprodione, isopamphos, isoprothiolane, kasugamycin, kresoxim-methyl (BAS 490F), mancopper, mancozeb, maneb, mepanipyrim, mepronil, mercuric chloride, mercuric oxide, mercurous chloride, metalaxyl, metam, metconazole, methasulfocarb, methyl isothiocyanate, metiram, myclobutanil, nabam, naphthenic acid, natamycin, neo-asozin (ferric methanearsonate), nickel bis(dimethyldithiocarbamate), nitrothal-isopropyl, nuarimol, octhilinone, ofurace, oxadixyl, oxine-copper, oxycarboxin,

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pefurazoate, penconazole, pencycuron, pentachlorophenol, phenylmercury acetate, 2-phenylphenol, phosdiphen, phthalide, piperalin, polyoxins, probenazole, prochloraz, procymidone, propamocarb, propiconazole, propineb, pyrazophos, pyrifenox, pyrimethanil, pyroquilon, quintozene, SSF-109, sulfur, tar oils, tebuconazole, tecnazene, tetraconazole, thiabendazole, thifluzamide, thiophanate-methyl, thiram, tolclofos-methyl, tolylfluanid, triadimenol, triazoxide, tricyclazole, tridemorph, triflumizole, triforine, triticonazole, validamycin, vinclozolin, zineb and ziram.

Suitable insecticides which may be incorporated in a composition with the compound of Formula I include: abamectin, acephate, azinphos-methyl, bifenthrin, buprofezin, carbofuran, chlorpyrifos, chlorpyrifos-methyl, cyfluthrin, beta-cyfluthrin, deltamethrin, diafenthiuron, diazinon, diflubenzuron, dimethoate, esfenvalerate, fenpropathrin, fenvalerate, fipronil, flucythrinate, tau-fluvalinate, fonophos, imidacloprid, isofenphos, malathion, metaldehyde, methamidophos, methidathion, methomyl, methoprene, methoxychlor, monocrotophos, oxamyl, parathion, parathion-methyl, permethrin, phorate, phosalone, phosmet, phosphamidon, pirimicarb, profenofos, rotenone, sulprofos, tebufenozide, tefluthrin, terbufos, tetrachlorvinphos, thiodicarb, tralomethrin, trichlorfon and triflumuron.

Suitable herbicides which may be incorporated in a composition with the compound of Formula I include: acetochlor, acifluorfen and its sodium salt, aclonifen, acrolein (2-propenal), alachlor, ametryn, amidosulfuron, amitrole, ammonium sulfamate, anilofos, asulam, atrazine, azimsulfuron, benazolin, benazolin, benfluralin, benfluralin, benfluresate, bensulfuron-methyl, bensulide, bentazone, bifenox, bromacil, bromoxynil, bromoxynil octanoate, butachlor, butralin, butylate, chlomethoxyfen, chloramben, chlorbromuron, chloridazon, chlorimuron-ethyl, chlornitrofen, chlorotoluron, chlorpropham, chlorsulfuron, chlorthal-dimethyl, cinmethylin, cinosulfuron, clethodim, clomazone, clopyralid, clopyralid-olamine, cyanazine, cycloate, cyclosulfamuron, 2,4-D and its butotyl, butyl, isoctyl and isopropyl esters and its dimethylammonium, diolamine and trolamine salts, daimuron, dalapon, dalapon-sodium, dazomet, 2,4-DB and its dimethylammonium, potassium and sodium salts, desmedipham, desmetryn, dicamba and its diglycolammonium, dimethylammonium, potassium and sodium salts, dichlobenil, dichlorprop, diclofop-methyl, 2-[4,5-dihydro-4-methyl-4-(1-methylethyl)-5-oxo-1H-imidazol-2-yl]-5-methyl-3pyridinecarboxylic acid (AC 263,222), difenzoquat metilsulfate, diflufenican, dimepiperate, dimethenamid, dimethylarsinic acid and its sodium salt, dinitramine, diphenamid, diquat dibromide, dithiopyr, diuron, DNOC, endothal, EPTC, esprocarb, ethalfluralin, ethametsulfuron-methyl, ethofumesate, ethyl \alpha,2-dichloro-5-[4-(difluoromethyl)-4,5-dihydro-3methyl-5-oxo-1H-1,2,4-triazol-1-yl]-4-fluorobenzenepropanoate (F8426), fenoxaprop-ethyl, fenoxaprop-P-ethyl, fenuron, fenuron-TCA, flamprop-methyl, flamprop-M-isopropyl, flamprop-M-methyl, flazasulfuron, fluazifop-butyl, fluazifop-P-butyl, fluchloralin, flumetsulam, flumiclorac-pentyl, flumioxazin, fluometuron, fluoroglycofen-ethyl, flupoxam, fluridone, flurochloridone, fluroxypyr, fomesafen, fosamine-ammonium, glufosinate, glufosinate-ammonium, glyphosate, glyphosate-isopropylammonium, glyphosate-sesquisodium, glyphosate-trimesium, halosulfuron-methyl, haloxyfop-etotyl, haloxyfop-methyl, hexazinone, imazamethabenz-methyl, imazamox (AC 299 263), imazapyr, imazaquin, imazaquin-ammonium, imazethapyr, imazethapyr-ammonium, imazosulfuron, ioxynil, ioxynil octanoate, ioxynil-sodium, isoproturon, isouron, isoxaben, isoxaflutole (RPA 201772), lactofen, lenacil, linuron, maleic hydrazide, MCPA

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and its dimethylammonium, potassium and sodium salts, MCPA-isoctyl, mecoprop, mecoprop-P, mefenacet, mefluidide, metam-sodium, methabenzthiazuron, methyl [[2-chloro-4-fluoro-5-[(tetrahydro-3-oxo-1H,3H-[1,3,4]thiadiazolo[3,4-a]pyridazin-1-ylidene)amino]phenyl]thioacetate (KIH 9201), methylarsonic acid and its calcium, monoammonium, monosodium and disodium salts, methyl [[[1-[5-[2-chloro-4-(trifluoromethyl)phenoxy]-2-nitrophenyl]-2methoxyethylidenelaminoloxylacetate (AKH-7088), methyl 5-[[[(4,6-dimethyl-2pyrimidinyl)amino]carbonyl]amino]sulfonyl]-1-(2-pyridinyl)-1H-pyrazole-4-carboxylate (NC-330), metobenzuron, metolachlor, metosulam, metoxuron, metribuzin, metsulfuron-methyl, molinate, monolinuron, napropamide, naptalam, neburon, nicosulfuron, norflurazon, oryzalin, oxadiazon, 3-oxetanyl 2-[[[(4,6-dimethyl-2-pyrimidinyl)amino]carbonyl]amino]sulfonyl]benzoate (CGA 277476), oxyfluorfen, paraquat dichloride, pebulate, pendimethalin, perfluidone, phenmedipham, picloram, picloram-potassium, pretilachlor, primisulfuron-methyl, prometon, prometryn, propachlor, propanil, propaquizafop, propazine, propham, propyzamide, prosulfuron, pyrazolynate, pyrazosulfuron-ethyl, pyridate, pyrithiobac, pyrithiobac-sodium, quinclorac, quizalofop-ethyl, quizalofop-P-ethyl, quizalofop-P-tefuryl, rimsulfuron, sethoxydim, siduron, simazine, sulcotrione (ICIA0051), sulfentrazone, sulfometuron-methyl, TCA, TCA-sodium, tebuthiuron, terbacil, terbuthylazine, terbutryn, thenylchlor, thiafluamide (BAY 11390), thifensulfuron-methyl, thiobencarb, tralkoxydim, tri-allate, triasulfuron, tribenuron-methyl, triclopyr, triclopyr-butotyl, triclopyr-triethylammonium, tridiphane, trifluralin, triflusulfuron-methyl, and vernolate.

The compositions containing the compound of Formula I may be used to control one or more of the pathogens listed in WO 90/12791 and others, including Alternaria cucumerina, Alternaria solani, Alternaria mali, Alternaria brassicola, Alternaria brassicae, Botryosphaeria obtusa, Bremia lectucae, Cercospora beticola, Cercospora arachidicola, Cercosporidium personatum, Cochliobolus sativus, Elsinoe ampelina, Elsinoe fawcetti, Fulvia fulva, Guignardia bidwelli, Gymnosporium juniperi-virginianae, Phomopsis viticola, Phytophthora nicotianae, Phytophthora megasperma, Pseudocercosporella capsallae, Puccinia asperagi, Puccinia hordei, Ramularia beticola, Rhynchosporium secalis, Septoria apiicola, Septoria lycopersici, Septoria tritici, Stemphilium solani, Peronophora tritici, Mycosphaerella musicola, Mycosphaerella fijiensis, Mycosphaerella citri, Mycosphaerella fragariae, Colletotrichum spp., Mucor spp, Penicillium spp., Gloeosporium spp., Glomerella spp., Gnomonia spp., Rhizopus spp., Aspergillus spp., Xanthomonas vesicatoria, Pseudomonas syringae and Helminthosporium spp.

The compound of Formula I or a composition containing it as already described may be applied by land or aerial application systems, mixed with soil, peat or other rooting media for the protection of plants against seed-borne, soil-borne or foliar pathogens.

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(74) Agents: HOUGHTON, Malcolm, John et al.; ICI Group Patents Services Dept., Shire Park, P.O. Box 6, Bessemer

(72) Inventors; and

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(54) Title: FUNGICIDAL COMPOSITION

$$\begin{array}{c|c}
N & CH_2 & C & R^2 \\
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(57) Abstract

A fungicidal composition comprising a carrier or diluent; a first active ingredient which is a compound of formula (I), and a second active ingredient which is a compound of formula (II), wherein R^1 is C_{1-4} alkyl or C_{3-6} cycloalkyl(C_{1-4}) alkyl, and R^2 is phenyl or phenyl (C1.4) alkyl; wherein the foregoing phenyl moieties are substituted with halogen; the relative amounts of the first and second active ingredients being such as to produce a synergistic effect.

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FUNGICIDAL COMPOSITION

The present invention relates to a fungicidal composition and to methods of using the composition to combat fungal infections of plants.

 (\underline{E}) -Methyl 2-[2-(6-(2-cyanophenoxy)pyrimidin-4-yloxy)phenyl]-3-methoxypropenoate, its use as a fungicide, compositions containing it, and methods of using it to combat fungal infections of plants are disclosed in EP-A2-0382375.

The present invention provides a fungicidal composition comprising a carrier or diluent; a first active ingredient which is a compound of formula (I), $[(\underline{E})$ -methyl 2-[2-(6-(2-cyanophenoxy)pyrimidin-4-yloxy)phenyl]-3-methoxypropenoate], and a second active ingredient which is a compound of formula (II) wherein R^1 is C_{1-4} alkyl or C_{3-6} cycloalkyl(C_{1-4})alkyl; and R^2 is phenyl or phenyl(C_{1-4})alkyl; wherein the foregoing phenyl moieties are substituted with halogen; the relative amounts of the first and second active ingredients being such as to produce a synergistic effect.

Alkyl and the alkyl moieties of C_{3-6} cycloalkyl(C_{1-4})alkyl and phenyl(C_{1-4})alkyl are in the form of straight or branched chains, and are, for example, methyl, ethyl, <u>n</u>-butyl or <u>tert</u>-butyl.

Cycloalkyl is, for example, cyclopropyl.

Halogen is preferably fluorine or chlorine.

The foregoing phenyl moieties are preferably 2- or 4- monosubstituted or 2,4- disubstituted.

In one aspect the present invention provides a fungicidal composition wherein the second active ingredient is a compound of formula (II) in which R^1 is a butyl group (especially <u>n</u>-butyl or <u>tert</u>-butyl) or 1-(cyclopropyl)eth-1-yl; and R^2 is mono- or di-chlorophenyl (especially 4-chlorophenyl or 2,4-dichlorophenyl) or monochlorophenyl(C_{1-2})alkyl (especially 2-chlorobenzyl or 2-(4-chlorophenyl)eth-1-yl).

In a further aspect the compound of formula (II) is a compound wherein R^1 is <u>n</u>-butyl and R^2 is 2,4-dichlorophenyl, or R^1 is <u>tert</u>-butyl and R^2 is 2-chlorobenzyl or 2-(4-chlorophenyl)eth-1-yl.

In another aspect there is provided a composition wherein the weight ratio of the first active ingredient to the second active ingredient is in the range from 400:1 to 10:90.

In a further aspect there is provided a fungicidal composition comprising a carrier or diluent; a first active ingredient which is a compound of formula (I), and a second active ingredient which is a compound

of formula (III); the relative amounts of the first and second active ingredients being such as to produce a synergistic effect, for example the weight ratio of the first active ingredient to the second active ingredient is in the range from 98:2 to 25:75, for example 9:1 to 25:75, and 98:2 to 9:1.

In a still further aspect there is provided a fungicidal composition comprising a carrier or diluent, a first active ingredient which is a compound of formula (I), and a second active ingredient which is a compound of formula (IV); the relative amounts of the first and second active ingredients being such as to produce a synergistic effect, for example the weight ratio of the first active ingredient to the second active ingredient is in the range from 400:1 to 10:90.

The compositions may be used to control one or more of the following pathogens: Pyricularia oryzae on rice and wheat and other Pyricularia spp. on other hosts.

<u>Puccinia recondita</u>, <u>Puccinia striiformis</u> and other rusts on wheat, <u>Puccinia hordei</u>, <u>Puccinia striiformis</u> and other rusts on barley, and rusts on other hosts e.g. turf, rye, coffee, pears, apples, peanuts, sugar beet, vegetables and ornamental plants.

Erysiphe graminis (powdery mildew) on barley, wheat, rye and turf and other powdery mildews on various hosts such as Sphaerotheca macularis on hops, Sphaerotheca fuliginea on cucurbits (e.g. cucumber), Podosphaera leucotricha on apple and Uncinula necator on vines.

Cochliobolus spp., Helminthosporium spp., Drechslera spp. (Pyrenophora spp.), Rhynchosporium spp., Septoria spp. (including Mycosphaerella graminicola and Leptosphaeria nodorum), Pseudocercosporella herpotrichoides and Gaeumannomyces graminis on cereals (e.g. wheat, barley, rye), turf and other hosts.

<u>Cercospora arachidicola</u> and <u>Cercosporidium personatum</u> on peanuts and other <u>Cercospora</u> species on other hosts, for example, sugar beet, bananas, soya beans and rice.

<u>Botrytis</u> <u>cinerea</u> (grey mould) on tomatoes, strawberries, vegetables, vines and other hosts and other Botrytis spp. on other hosts.

Alternaria spp. on vegetables (e.g. cucumber), oil-seed rape, apples, tomatoes, cereals (e.g. wheat) and other hosts.

<u>Venturia</u> spp. (including <u>Venturia inaequalis</u> (scab)) on apples, pears, stone fruit, tree nuts and other hosts.

Cladosporium spp. on a range of hosts including cereals (e.g. wheat).

Monilinia spp. on stone fruit, tree nuts and other hosts.

Didymella spp. on tomatoes, turf, wheat and other hosts.

Phoma spp. on oil-seed rape, turf, rice, potatoes, wheat and other
hosts.

Aspergillus spp. and Aureobasidium spp. on wheat, lumber and other hosts.

Ascochyta spp. on peas, wheat, barley and other hosts.

<u>Plasmopara viticola</u> on vines. Other downy mildews such as <u>Bremia lactucae</u> on lettuce, <u>Peronospora</u> spp. on soybeans, tobacco, onions and other hosts, <u>Pseudoperonospora humuli</u> on hops and <u>Pseudoperonospora</u> cubensis on cucurbits.

Pythium spp. on turf and other hosts.

<u>Phytophthora</u> <u>infestans</u> on potatoes and tomatoes and other <u>Phytophthora</u> spp. on vegetables, strawberries, avocado, pepper, ornamentals, tobacco, cocoa and other hosts.

Thanatephorus <u>cucumeris</u> on rice and turf and other <u>Rhizoctonia</u> species on various hosts such as wheat and barley, vegetables, cotton and turf.

Sclerotinia spp. on turf, peanuts, oil-seed rape and other hosts.

Sclerotium spp. on turf, peanuts and other hosts.

<u>Colletotrichum</u> spp. on a range of hosts including turf, coffee and vegetables.

Laetisaria fuciformis on turf.

Mycosphaerella spp. on banana, peanut, citrus, pecan, papaya and other hosts.

Diaporthe spp. on citrus, soybean, melon, pear, lupin and other hosts.

Elsinoe spp. on citrus, vines, olives, pecans, roses and other hosts.

Pyrenopeziza spp. on oil-seed rape and other hosts.

Oncobasidium theobromae on cocoa causing vascular streak dieback.

<u>Fusarium spp., Typhula spp., Microdochium nivale, Ustilago spp.,</u>
<u>Urocystis spp., Tilletia spp., and Claviceps purpurea</u> on a variety of hosts but particularly wheat, barley, turf and maize.

<u>Verticillium</u> spp. on a range of hosts including cotton, potatoes, tomatoes and hops.

Ramularia spp. on sugar beet and other hosts.

Post-harvest diseases particularly of fruit (e.g. <u>Pencillium digitatum</u> and P. <u>italicum and Trichoderma viride</u> on oranges, <u>Colletotrichum musae</u> and <u>Gloeosporium musarum</u> on bananas and <u>Botrytis cinerea</u> on grapes).

Other pathogens on vines, notably Eutypa lata, Guignardia bidwellii,

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<u>Phellinus igniarus</u>, <u>Phomopsis viticola</u>, <u>Pseudopezicula tracheiphila</u> and Stereum hirsutum.

Other pathogens on cereals, notably <u>Selonophoma donacis</u>, <u>Sclerophthora</u> spp., <u>Sclerospora sorghi</u>, <u>Nigrospora oryzae</u>, <u>Trichometasphaeria turcica</u>, <u>Cephalosporium gramineum</u>, <u>Epicoccum spp.</u>, <u>Stemphylium spp.</u>, <u>Sporobolomyces spp.</u>, <u>Cryptosporium spp.</u>, <u>Dilophospora alopecuri</u>, <u>Phaeoseptoria urvilleana</u>, <u>Phyllachora graminis</u>, <u>Omphalina pixidata and Platyspora pentamera</u>.

Other pathogens on turf, notably <u>Agaricus</u> spp., <u>Coprinus</u>

<u>psychromorbidus</u>, <u>Epichloe typhina</u>, <u>Lepiota spp.</u>, <u>Leptosphaeria korrae</u>,

<u>Lycoperdon spp.</u>, <u>Magnaporthe poae</u>, <u>Marasmius oreades</u>, <u>Myriosclerotinia</u>

borealis, Nigrospora sphaerica, <u>Phialophora spp.</u> and <u>Pyricularia grisea</u>.

Other pathogens on lumber, notably <u>Cephaloascus fragrans</u>, <u>Ceratocystis</u> spp., <u>Chaetomium globosum</u>, <u>Coniophora puteana</u>, <u>Coriulus veriscolor</u>, <u>Dothiorella spp., <u>Gliocladium virens</u>, <u>Gloeophyllum trabeum</u>, <u>Ophiostoma piceae</u>, <u>Penicillium spp.</u>, <u>Phialophora spp.</u>, <u>Phlebia gigantea</u>, <u>Poria placenta</u>, <u>Rhinocladiella atrovirens</u>, <u>Schizophyllum vaillanti</u>, <u>Sclerophoma pityophila</u>, <u>Sistotrema brinkmannii</u>, <u>Sphaeropsis sapina</u>, <u>Trametes veriscolor</u>, <u>Trichoderma pseudokoningii</u>, <u>Trichoderna viride</u> and slime mould.</u>

Fungal vectors of viral diseases e.g. <u>Polymyxa graminis</u> on cereals as the vector of barley yellow mosaic virus (BYMV).

Some of the compositions show a broad range of activities against fungi $\underline{\text{in }}$ $\underline{\text{vitro}}.$

The active ingredients may move acropetally/locally in plant tissue.

The invention therefore provides a method of combating fungi which comprises applying to a plant, to a seed of a plant or to the locus of the plant or seed a fungicidally effective amount of a composition as hereinbefore defined.

It is preferred that all compositions, both solid and liquid formulations, comprise 0.0001 to 95%, more preferably 1 to 85%, for example 1 to 25% or 25 to 60%, of the first and second active ingredients combined.

When applied the foliage of plants, the compounds of the invention are applied at rates of 0.1g to 10Kg, preferably 10g to 8Kg, more preferably 10g to 4Kg, of the first and second active ingredients combined per hectare.

When used as seed dressings, the compositions of the invention are used at rates of 0.0001g (for example 0.001g or 0.05g) to 10g, preferably 0.1g to 8g, more preferably 0.1g to 4g, of the first and second active ingredients combined per kilogram of seed.

The compositions can be applied in a number of ways. For example, they can be applied directly to the foliage of a plant, to seeds or to other medium in which plants are growing or are to be planted, or they can be sprayed on, dusted on or applied as a cream or paste formulation, or they can be applied as a vapour or as slow release granules.

Application can be to any part of the plant including the foliage, stems, branches or roots, or to soil surrounding the roots, or to the seed before it is planted, or to the soil generally, to paddy water, to irrigation water or to hydroponic culture systems. The invention compositions may also be injected into plants or sprayed onto vegetation using electrodynamic spraying techniques or other low volume methods.

The term "plant" as used herein includes seedlings, bushes and trees. Furthermore, the fungicidal method of the invention includes preventative, protectant, prophylactic and eradicant treatments.

The physical nature of the composition used in any instance will depend upon the particular purpose envisaged.

The compositions may be in the form of dustable powders or granules comprising the first and second active ingredients and a solid diluent or carrier, for example, fillers such as kaolin, bentonite, kieselguhr, dolomite, calcium carbonate, talc, powdered magnesia, fuller's earth, gypsum, diatomaceous earth and china clay. Such granules can be preformed granules suitable for application to the soil without further treatment. These granules can be made either by impregnating pellets of filler with the active ingredient or by pelleting a mixture of the active ingredient and powdered filler. Compositions for dressing seed may include an agent (for example, a mineral oil) for assisting the adhesion of the composition to the seed; alternatively the active ingredient can be formulated for seed dressing purposes using an organic solvent (for example, N-methylpyrrolidone, propylene glycol or N,N-dimethylformamide). The compositions may also be in the form of wettable powders or water dispersible granules comprising wetting or dispersing agents to facilitate the dispersion in liquids. The powders and granules may also contain fillers and suspending agents.

Compositions for pelleting seed comprise the first and second active ingredients, a solid carrier (such as clay or chalk) and a binding agent (such as a cellulosic or polysaccharide material).

The compositions may also be in the form of soluble powders or granules, or in the form of solutions in polar solvents.

Soluble powders may be prepared by mixing the active ingredient with a water-soluble salt such as sodium bicarbonate, sodium carbonate, magnesium sulphate or a polysaccharide, and a wetting or dispersing agent to improve water dispersibility/solubility. The mixture may then be ground to a fine powder. Similar compositions may also be granulated to form water-soluble granules. Solutions may be prepared by dissolving the active ingredient in polar solvents such as ketones, alcohols and glycol ethers. These solutions may contain surface active agents to improve water dilution and prevent crystallisation in a spray tank.

Emulsifiable concentrates or emulsions may be prepared by dissolving the active ingredient in an organic solvent optionally containing a wetting or emulsifying agent and then adding the mixture to water which may also contain a wetting or emulsifying agent. Suitable organic solvents are aromatic solvents such as alkylbenzenes and alkylnaphthalenes, ketones such as cyclohexanone and methylcyclohexanone, chlorinated hydrocarbons such as chlorobenzene and trichlorethane, and alcohols such as benzyl alcohol, furfuryl alcohol, butanol and glycol ethers.

Suspension concentrates of largely insoluble solids may be prepared by ball or bead milling with a dispersing agent with a suspending agent included to stop the solid settling.

Compositions to be used as sprays may be in the form of aerosols wherein the formulation is held in a container under pressure of a propellant, e.g. fluorotrichloromethane or dichlorodifluoromethane.

The first and second active ingredients can be mixed in the dry state with a pyrotechnic mixture to form a composition suitable for generating in enclosed spaces a smoke containing the active ingredients.

Alternatively, the compositions may be used in micro-encapsulated form. They may also be formulated in biodegradable polymeric formulations to obtain a slow, controlled release of the active substance.

By including suitable additives, for example additives for improving the uptake, distribution, adhesive power and resistance to rain on treated surfaces, the different compositions can be better adapted for various utilities. Other additives may be included to improve the biological efficacy of the various formulations. Such additives can be surface active materials to improve the wetting and retention on surfaces treated with the formulation and also the uptake and mobility of the active material, or additionally can include oil based spray additives, for example, certain mineral oil and natural plant oil (such as soya bean and rape seed oil)

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additives, or blends of them with other adjuvants, have been found to enhance several-fold foliar activity against, for example, <u>Puccinia hordei</u>.

The invention compositions can be used as mixtures with fertilisers (e.g. nitrogen-, potassium- or phosphorus-containing fertilisers). Granules comprising only pieces of fertiliser incorporating, for example coated with, the composition are preferred. Such granules suitably contain up to 25% by weight of the first and second active ingredients. The invention therefore also provides a fertiliser and a composition of the invention.

Wettable powders, emulsifiable concentrates and suspension concentrates will normally contain surfactants, e.g. a wetting agent, dispersing agent, emulsifying agent or suspending agent. These agents can be cationic, anionic or non-ionic agents.

Suitable cationic agents are quaternary ammonium compounds, for example, cetyltrimethylammonium bromide. Suitable anionic agents are soaps, salts of aliphatic monoesters of sulphuric acid (for example, sodium lauryl sulphate), and salts of sulphonated aromatic compounds (for example, sodium dodecylbenzenesulphonate, sodium, calcium or ammonium lignosulphonate, butylnaphthalene sulphonate, and a mixture of sodium diisopropyl- and triisopropylnaphthalene sulphonates).

Suitable non-ionic agents are the condensation products of ethylene oxide with fatty alcohols such as oleyl or cetyl alcohol, or with alkyl phenols such as octyl— or nonylphenol and octylcresol. Other non-ionic agents are the partial esters derived from long chain fatty acids and hexitol anhydrides, the condensation products of the said partial esters with ethylene oxide, and the lecithins. Suitable suspending agents are hydrophilic colloids (for example, polyvinylpyrrolidone and sodium carboxymethylcellulose), and swelling clays such as bentonite or attapulgite.

Compositions for use as aqueous dispersions or emulsions are generally supplied in the form of a concentrate containing a high proportion of the active ingredient, the concentrate being diluted with water before use. These concentrates should preferably be able to withstand storage for prolonged periods and after such storage be capable of dilution with water in order to form aqueous preparations which remain homogeneous for a sufficient time to enable them to be applied by conventional spray equipment. The concentrates may conveniently contain up to 95%, suitably 1-85%, for example 1-25% or 25-60%, by weight of the active ingredients.

After dilution to form aqueous preparations, such preparations may contain varying amounts of the active ingredient depending upon the intended purpose, but an aqueous preparation containing 0.0001 to 10%, for example 0.005 to 10%, by weight of active ingredient may be used.

The compositions of this invention may contain other compounds having biological activity, e.g. compounds having similar or complementary fungicidal activity or which possess plant growth regulating, herbicidal or insecticidal activity.

An additional fungicidal compound may be present in the composition of the invention. By including another fungicide, the resulting composition can have a broader spectrum of activity or a greater level of intrinsic activity than the composition of the invention alone. Examples of fungicidal compounds which may be included in the composition of the invention are (±)-cis-1-(4-chlorophenyl)-2-(1H-1,2,-triazol-1-yl)--cycloheptanol, (2RS, 3RS)-1-[3-(2-chlorophenyl)-2-(4-fluorophenyl)--oxiran-2-ylmethyl]-1H-1,2,4-triazole, (RS)-1-aminopropylphosphonic acid, (RS)-4-(4-chlorophenyl)-2-phenyl-2-(1H-1,2,4-triazol-1-ylmethyl)butyronitrile, (Z)-N-but-2-enyloxymethyl-2-chloro-2',6'-diethylacetanilide, 1-(2-cyano-2-methoxyiminoacety1)-3-ethyl urea, 3-(2,4-dichloropheny1)-2--(1H-1,2,4-triazol-1-yl)quinazolin-4(3H)-one, 4-(2,2-difluoro-1,3--benzodioxol-4-yl)pyrrole-3-carbonitrile, 4-bromo-2-cyano-N,N-dimethyl-6--trifluoromethylbenzimidazole-1-sulphonamide, 5-ethyl-5,8-dihydro-8-oxo--(1,3)-dioxol-(4,5-g)quinoline-7-carboxylic acid, α -[N-(3-chloro-2,6--xylyl)-2-methoxyacetamido]-γ-butyrolactone, alanycarb, aldimorph, ampropylfos, anilazine, azaconazole, BAS 490F, benalaxyl, benomyl, biloxazol, binapacryl, bitertanol, blasticidin S, bromuconazole, bupirimate, butenachlor, buthiobate, captafol, captan, carbendazim, carbendazim chlorhydrate, carboxin, chinomethionate, chlorbenzthiazone, chloroneb, chlorothalonil, chlorozolinate, clozylacon, copper containing compounds such as copper oxychloride, copper oxyquinolate, copper sulphate and Bordeaux mixture, cycloheximide, cymoxanil, cyproconazole, cyprofuram, di-2-pyridyl disulphide 1,1'-dioxide, dichlofluanid, dichlone, diclobutrazol, diclomezine, dicloran, didecyl dimethyl ammonium chloride, diethofencarb, difenoconazole, 0,0-di-iso-propyl-S-benzyl thiophosphate, dimefluazole, dimetconazole, dimethomorph, dimethirimol, diniconazole, dinocap, dipyrithione, ditalimfos, dithianon, dodemorph, dodine, doguadine, edifenphos, epoxiconazole, etaconazole, ethirimol, ethoxyquin, ethyl (Z)-N-benzyl-N-([methyl(methyl-thioethylideneamino-oxycarbonyl)amino|thio)-

 $-\beta$ -alaninate, etridiazole, fenaminosulph, fenapanil, fenarimol, fenbuconazole, fenfuram, fenpiclonil, fenpropidin, fenpropimorph, fentin acetate, fentin hydroxide, ferbam, ferimzone, fluazinam, fluoroimide, fluotrimazole, flutolanil, flutriafol, flusilazole, folpet, fuberidazole, furalaxyl, furconazole-cis, guazatine, hydroxyisoxazole, hymexazole, imazalil, imibenconazole, ipconazole, iprobenfos, iprodione, isopropanyl butyl carbamate, isoprothiolane, kasugamycin, mancozeb, maneb, mepanipyrim, mepronil, metalaxyl, metconazole, methfuroxam, metiram, metiram-zinc, metsulfovax, myclobutanil, neoasozin, nickel dimethyldithiocarbamate, nitrothal-isopropyl, nuarimol, ofurace, organomercury compounds, oxadixyl, oxolinic acid, oxycarboxin, pefurazoate, penconazole, pencycuron, phenazin oxide, phosetyl-Al, phosphorus acids, phthalide, polyoxin D, polyram, probenazole, prochloraz, procymidone, propamocarb, propamocarb hydrochloride, propiconazole, propineb, propionic acid, prothiocarb, pyracarbolid, pyrazophos, pyrifenox, pyroquilon, pyroxyfur, pyrrolnitrin, quaternary ammonium compounds, quinconazole, quinomethionate, quintozene, rabenazole, sodium pentachlorophenate, SSF 126, streptomycin, sulphur, tebuconazole, techlofthalam, tecnazene, tetraconazole, thiabendazole, thicarbanil, thicyofen, 2-(thiocyanomethylthio)benzothiazole thiophanate-methyl, thiram, timibenconazole, tolclofos-methyl, tolylfluanid, triacetate salt of 1,1'-iminodi-(octamethylene)diguanidine, triadimefon, triadimenol, triazbutyl, triazoxide, tricyclazole, tridemorph, triforine, triflumizole, triticonazole, validamycin A, vapam, vinclozolin, zineb and ziram. The compounds of general formula (I) can be mixed with soil, peat or other rooting media for the protection of plants against seed-borne, soil-borne or foliar fungal diseases.

Suitable insecticides which may be incorporated in a composition of the present invention include natural pyrethrins or pyrethroids such as permethrin, fenvalerate, deltamethrin, cyhalothrin, lambda-cyhalothrin, cypermethrin, α - and β -cypermethrin, cycloprothrin, tefluthrin, empenthrin, ethofenprox, tetramethrin, bioallethrin, fenfluthrin, prallethrin, 5-benzyl-3-furylmethyl (E)-(1R,3S)-2,2-dimethyl-3-(2-oxothiolan-3-yl-idenemethyl)cyclopropane carboxylate and pentafluorobenzyl (cis)-3-[2-fluoro-2-(methoxycarbonyl)ethenyl]-2,2-dimethylcyclopropane carboxylate; organophosphates such as profenofos, propaphos, sulprofos, dichlorvos, methyl parathion, azinphos-methyl, demeton-s-methyl, heptenophos, thiometon, fenamiphos, monocrotophos, triazophos, methamidophos, dimethoate, phenthoate, phosphamidon, malathion,

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chlorpyrifos, phosalone, fensulphothion, fenthion, formothion, isoxathion, fonofos, phorate, phoxim, pyrimiphos-methyl, fenitrothion and diazinon; carbamates (including aryl carbamates) such as pirimicarb, cloethocarb, carbofuran, carbonyl, isoprocarb, ethiofencarb, aldicarb, thiofurox, carbosulfan, bendiocarb, fenobucarb, propoxur, oxamyl and XMC; benzoyl ureas such as triflumuron and chlorofluazuron; organic tin compounds such as cyhexatin, fenbutatin oxide and azocyclotin; macrolides such as avermectins or milbemycins, for example abamectin, avermectin and milbemycin; hormones and synthetic mimics thereof such as juvenile hormone, juvabione, ecdysones, methoprene and hydroprene; pheromones; and organochlorine compounds such as benzene hexachloride, DDT, chlordane, dieldrin and endosulfan.

In addition to the major chemical classes of insecticide listed above, other insecticides having particular targets may be employed in the mixture an appropriate. For instance selective insecticides for particular crops, for example stem borer specific insecticides for use in rice such as cartap or buprofezin, can be employed. Alternatively, insecticides or acaricides specific for the control of specific insect growth stages, for example ovolarvicides such as clofentezine, amitraz, chlordimeform, flubenzimine, hexythiazox and tetradifon; motilicides such as dicofol or propargite; adulticides such as bromopropylate, chlorobenzilate; or insect growth regulators such as hydramethylnon, cyromazine, methoprene, chlorfluazuron and diflubenzuron may also be included in the compositions.

Examples of suitable insecticide synergists for use in the compositions include piperonyl butoxide, sesamex and dodecyl imidazole.

Plant growth regulating compounds are compounds which control weeds or seedhead, formation, or selectively control the growth of less desirable plants (e.g. grasses).

Examples of suitable plant growth regulating compounds for use in the compositions of the present invention are 1-(4-chlorophenyl)-4,6-di-methyl-2-oxo-1,2-dihydropyridine-3-carboxylic acid; methyl-3,6-dichloroanisate; abscisic acid; daminozide; difenzoquat; dikegulac; ethephon; fenpentezol; fluoridamid; inabenfide; isopyrimol; long chain fatty alcohols and acids; maleic hydrazide; mefluidide; fenchlorazole-ethyl; chloine chloride; ethephon; morphactins (e.g. chlorfluoroecol); paclobutrazol; substituted benzoic acid (e.g. triiodobenzoic acid); substituted quaternary ammonium and phosphonium compounds (e.g. chloromequat, chlorphonium or mepiquat chloride); tecnazene; the auxins (e.g. indoleacetic acid, indolebutyric

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acid, naphthylacetic acid or naphthoxyacetic acid); the cytokinins (e.g. benzimidazole, benzyladenine, benzylaminopurine, diphenylurea or kinetin); the gibberellins (e.g. GA_3 , GA_4 or GA_7); triapenthenol; benzo-2,1,3-thiadiazin-4-one-2,2-dioxides such as bentazone; hormone herbicides, particularly the phenoxy alkanoic acids such as MCPA, MCPA-thioethyl, dichlorprop, 2,4,5-T, MCPB, 2,4-D, 2,4-DB, mecoprop, trichlopyr, fluroxypyr, clopyralid, and their derivatives (eg. salts, esters and amides); 1,3 dimethylpyrazole derivatives such as pyrazoxyfen, pyrazolate and benzofenap; Dinitrophenols and their derivatives (eg. acetates) such as dinoterb, dinoseb and its ester, dinoseb acetate; dinitroaniline herbicides such as dinitramine, trifluralin, ethalflurolin, pendimethalin, oryzalin; arylurea herbicides such as diuron, flumeturon, metoxuron, neburon, isoproturon, chlorotoluron, chloroxuron, linuron, monolinuron, chlorobromuron, daimuron, methabenzthiazuron; phenylcarbamoyloxyphenylcarbamates such as phenmedipham and desmedipham; 2-phenylpyridazin-3-ones such as chloridazon and norflurazon; uracil herbicides such as lenacil, bromacil and terbacil; triazine herbicides such as atrazine, simazine, aziprotryne, cyanazine, prometryn, dimethametryn, simetryne, and terbutryn; phosphorothioate herbicides such as piperophos, bensulide, and butamifos; thiolcarbamate herbicides such as cycloate, vernolate, molinate, thiobencarb, butylate*, EPTC*, tri-allate, di-allate, esprocarb, thiocarbazil, pyridate, and dimepiperate; 1,2,4-triazin-5-one herbicides such as metamitron and metribuzin; benzoic acid herbicides such as 2,3,6-TBA, dicamba and chloramben; anilide herbicides such as pretilachlor, butachlor, alachlor, propachlor, propanil, metazachlor, metolachlor, acetochlor, and dimethachlor; dihalobenzonitrile herbicides such as dichlobenil, bromoxynil and ioxynil; haloalkanoic herbicides such as dalapon, TCA and salts thereof; diphenylether herbicides such as lactofen, fluroglycofen or salts or ester thereof, nitrofen, bifenox, aciflurofen and salts and esters thereof, oxyfluorfen, fomesafen, chlornitrofen and chlomethoxyfen; phenoxyphenoxypropionate herbicides such as diclofop and esters thereof such as the methyl ester, fluazifop and esters thereof, benzoylprop and esters thereof, haloxyfop and esters thereof, quizalofop and esters thereof and fenoxaprop and esters thereof such as the ethyl ester; cyclohexanedione herbicides such as alloxydim and salts thereof, sethoxydim, cycloxyidim, tralkoxydim, and clethodim; sulfonyl urea herbicides such as chlorosulfuron, sulfometuron, metsulfuron and esters thereof; benzsulfuron and esters thereof such as DPX-M6313,

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chlorimuron and esters such as the ethyl ester thereof pirimisulfuron and esters such as the methyl ester thereof, 2-[3-(4-methoxy-6-methyl-1,3,5--triazin-zyl)-3-methylureido-sulphonyl) benzoic acid esters such as the methyl ester thereof (DPX-LS300) and pyrazosulfuron; imidazolidinone herbicides such as imazaquin, imazamethabenz, imazapyr and isopropylammonium salts thereof, imazethapyr; arylanilide herbicides such as flamprop and esters thereof, benzoylprop-ethyl, diflufenican; amino acid herbicides such as glyphosate and glufosinate and their salts and esters, sulphosate and bialaphos; organoarsenical herbicides such as monosodium methanearsonate (MSMA); herbicidal amide derivative such as napropamide, propyzamide, asulam, carbetamide, tebutam, bromobutide, isoxaben, naproanilide and naptalam; miscellaneous herbicides including ethofumesate, cinmethylin, difenzoquat and salts thereof such as the methyl sulphate salt, clomazone, oxadiazon, bromofenoxim, barban, tridiphane, flurochloridone, quinclorac, dithiopyr and mefanacet; and contact herbicides (such as bipyridylium herbicides for example those in which the active entity is paraquat and those in which the active entity is diquat), and mixtures of any of the foregoing. (* These compounds are preferably employed in combination with a safener such as dichlormid.)

The following Examples illustrate the invention.

EXAMPLE 1

Testing of a composition of the invention against <u>Puccinia</u> <u>hordei</u> (barley brown rust) in a field test on barley in test area $20m^2$ (four replicates).

The compounds were applied as suspension concentrate formulations with a tank-mixed adjuvant and were diluted with water to the required concentration.

The compounds were applied using knapsack sprayers in a vater volume of 300 l/ha. Two applications were made between GSZ 30 and GSZ 65 depending upon disease pressure and local recommendations (GSZ = Zadoks Growth Stage). Disease control was assessed as percentage brown rust on specific leaves by visual estimation of 25 tillers per plot.

Results are expressed as mean absolute percentage disease levels, and untreated plots had a mean absolute disease level of 80.40%.

Application rate g ai/ha		Mean absolute disease level		
Compound (I)	Compound (IV)	(on leaf 2; 15 days after application 2)		
25	0	33.51%		
0	31.25	26.76%		
25	31.25	6.65%		

EXAMPLE 2

Testing of a composition of the invention against <u>Venturia inaequalis</u> (apple scab).

Young apple seedlings were grown in 3.8cm (1.5") diameter plastic pots containing John Innes Potting Compost (No. 1).

The test compounds, formulated as suspension concentrates or a solution in acetone, were diluted in water to the required concentration immediately before application.

The test compounds were applied to the underside of the apple seedling leaves either one day before (protectant) or three days after (eradicant) a suspension of <u>Venturia inaequalis</u> spores was applied to the same surface.

After inoculation, the plants were put into an appropriate environment to allow infection to proceed and then incubated until the disease was ready for assessment (10-12 days after inoculation). <u>Venturia inaequalis</u> infection was assessed as the percentage area of sporulating disease on a leaf in comparison with the untreated control. Four replicate plants were assessed per treatment.

Eradicant Treatment

Application rate ppm ai		Percentage area of sporulating		
Compound (I)	Compound (III)	disease		
0.2	0	70.7		
0	0.2	76.9		
0.2	0.2	14.1		

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Protectant Treatment

Application	n rate ppm ai	Percentage area of sporulating
Compound (I)	Compound (III)	disease
0.2	0	74.5
0	0.2	109
0.2	0.2	20.4

These Examples give data for the two components present in the compositions. By using Limpel's formula, these data demonstrate that the observed activity of the compositions comprising two active ingredients is greater than would be expected.

Limpel's formula (Pesticide Science (1987) 19 309-315 at 312) is:

$$E = X+Y - \frac{XY}{100}$$

in which

E is the expected fungicidal activity of substances A+B at p+q Kg/ha;

X is the percentage fungicidal activity of substance A at a rate of p kg active ingredient/ha; and

Y is the percentage fungicidal activity of substance B at a rate of q kg active ingredient/ha.

If the fungicidal activity observed is greater than the value E calculated according to Limpel, then the combination of A and B is synergistic.

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CHEMICAL FORMULAE

(IN DESCRIPTION)

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BNSDOCID: <WO__8322921A1__>

CLAIMS

 A fungicidal composition comprising a carrier or diluent; a first active ingredient which is a compound of formula (I):

and a second active ingredient which is a compound of formula (II):

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wherein R^1 is C_{1-4} alkyl or C_{3-6} cycloalkyl(C_{1-4})alkyl; and R^2 is phenyl or phenyl(C_{1-4})alkyl; wherein the foregoing phenyl moieties are substituted with halogen; the relative amounts of the first and second active ingredients being such as to produce a synergistic effect.

2. A fungicidal composition as claimed in claim 1 wherein the second active ingredient is a compound of formula (III) or (IV).

$$\begin{array}{c|c}
 & C1 \\
 & N \\
 & CH_2 \\
 & CCH_3 \\
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- 3. A method of combating fungal infections of plants comprising applying to the plant, to the seed of the plant or to the locus of the plant or seed a composition as claimed in claim 1.
- 4. A method of combating fungal infections of plants caused by <u>Puccinia</u> spp, the method comprising applying to the plant, to the seed of the plant or to the locus of the plant or seed a fungicidal composition comprising a diluent or carrier; a first active ingredient which is a compound of formula (I); and a second active ingredient which is a compound of formula (IV); the relative amounts of the first and second active ingredients being such as to produce a synergistic effect.
- 5. A method of combating fungal infections of plants caused by <u>Venturia</u> spp, the method comprising applying to the plant, to the seed of the plant or to the locus of the seed or plant a fungicidal composition comprising a diluent or carrier; a first active ingredient which is a compound of formula (I); and a second active ingredient which is a compound of formula (III); the relative amounts of the first and second active ingredients being such as to produce a synergistic effect.
- 6. A composition as claimed in claim 1 or 2 or a composition as used in claim 4 or 5 wherein the weight ratio of the first active ingredient to the second active ingredient is in the range from 400:1 to 10:90.
- 7. A method of combating fungal infections of plants comprising applying to a plant, to the seed of a plant or to the locus of a seed or plant a fungicidally effective amount of a composition as claimed in claim 1.

International A tion No I. CLASSIFICATION OF SUBJEC TER (if several classification symbols apply, indicate According to International Patent Classification (IPC) or to both National Classification and IPC Int.Cl. 5 A01N43/653; //(A01N43/653,43:54) II. FIELDS SEARCHED Minimum Documentation Searched? Classification Symbols Classification System **A01N** Int.Cl. 5 Documentation Searched other than Minimum Documentation to the Extent that such Documents are Included in the Fields Searched III. DOCUMENTS CONSIDERED TO BE RELEVANT? Category o Citation of Document, 11 with indication, where appropriate, of the relevant passages 12 Relevant to Claim No.13 X EP,A,O 382 375 (IMPERIAL CHEMICAL 1-5.7INDUSTRIES) 16 August 1990 cited in the application see page 3, line 1 - line 37 see page 5; table 1; compound no. 9 see page 29, line 51 - page 30, line 26 P,X RESEARCH DISCLOSURE 1-7 no. 338, 10 June 1992, HAVANT GB pages 506 - 510 ANONYMOUS 'Mixtures of Fungicides and Insecticides' disclosure no. 33893 o Special categories of cited documents: 10 "T" later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the "A" document defining the general state of the art which is not considered to be of particular relevance earlier document but published on or after the international "X" document of particular relevance; the claimed invention filing date annot be considered novel or cannot be considered to document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another involve an inventive step "Y" document of particular relevance; the claimed invention citation or other special reason (as specified) cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled "O" document referring to an oral disclosure, use, exhibition or other means document published prior to the international filing date but later than the priority date claimed "A" document member of the same patent family IV. CERTIFICATION Date of the Actual Completion of the International Search Date of Mailing of this International Search Report 27 JULY 1993 0 6. 08. 93 International Searching Authority Signature of Authorized Officer LAMERS W. **EUROPEAN PATENT OFFICE**

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III. DOCTIN	TENTS CONSIDERED TO BE RELL (CONTINUED FROM THE SECOND SHEE	
Category °	Citation of Document, with indication, where appropriate, of the relevant passages	Relevant to Claim No.
	The state of the s	
Р,Х	DATABASE WPIL Week 9313, Derwent Publications Ltd., London, GB; AN 93-107493 see abstract & RESEARCH DISCLOSURE no. 346, 10 February 1993, HAVANT GB ANONYMOUS 'Fungicidal mixts contg. E-me thyl(((cyano:phenoxy)pyrimidinyl:oxy)pheny 1)methoxy:propenoate and further fungicides' disclosure no. 346121	1,2,7
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	·	

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27/07/93

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EP-A-0382375	16-08-90	AU-B- AU-A- CN-A- US-A-	625501 4905290 1047286 5145856	16-07-92 16-08-90 28-11-90 08-09-92	
EP-A-0415569	06-03-91	AU-A- CN-A- JP-A-	6022090 1049771 3193705	07-03-91 13-03-91 23-08-91	

For more details about this annex : see Official Journal of the European Patent Office, No. 12/82

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Before the expiration of the time limit for amending the claims and to be republished in the event of the receipt of amendments.

(54) Title: FUNGICIDAL MIXTURES

(57) Abstract

Advantageous combinations are disclosed comprising (1) at least one compound selected from the quinazolinones of Formula (I), N-oxides and agriculturally suitable salts thereof and (2) at least one compound selected from: (a) compounds acting at the bc_I complex of the fungal mitochondrial respiratory electron transfer site and (b) compounds that control fungal disease by inhibiting sterol biosynthesis. D is O or S and R^1 - R^4 are as defined in the disclosure. Included are fungicidal

$$R^3$$
 N
 R^1
 OR^2

compositions comprising fungicidally effective amounts of the compound combinations of the invention and a method for controlling plant diseases caused by fungal plant pathogens comprising applying to the plant or portion thereof, or to the plant seed or seedling, a fungicidally effective amount of the compound combinations of the invention.

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TITLE

FUNGICIDAL MIXTURES BACKGROUND OF THE INVENTION

Fungicides that effectively control plant diseases are in constant demand by growers. Plant diseases are highly destructive, difficult to control and quickly develop resistance to commercial fungicides. Combinations of pesticides are often used to facilitate disease control, to broaden spectrum of control and to retard resistance development. It is recognized in the art that the advantages of particular pesticide combinations can often vary, depending on such factors as the particular plant and plant disease to be treated, and the treatment conditions. Accordingly, there is an ongoing search for advantageous combinations of pesticides.

International Patent Application WO 94/26722 discloses certain quinazolinone compounds as fungicides (e.g., 6-bromo-3-propyl-2-propyloxy-4(3H)-quinazolinone, 6.8-diiodo-3-propyl-2-propyloxy-4(3H)-quinazolinone and 6-iodo-3-propyl-2propyloxy-4(3H)-quinazolinone). International Patent Application WO 95/14009 discloses certain triazolone fungicides (e.g., 2,4-dihydro-5-methoxy-2-methyl-4-[2-[[[1-[3-(trifluoromethyl)phenyl]ethylidene]amino]oxy]-methyl]phenyl]-3H-1,2,4triazol-3-one). Kresoxim-methyl (BAS 490F) is a fungicide under consideration for the control of plant diseases, especially fungal diseases of wheat, grapes and fruits. (See E. Ammermann, G. Lorenz, K. Schelberger, B. Wenderoth, H. Sauter and C. Rentzea "BAS 490F - A Broad-Spectrum Fungicide with a New Mode of Action" in Brighton Crop Protection Conference - Pests and Diseases - 1992, 1, 403-410.) Azoxystrobin (ICIA5504) is a fungicide under consideration for the control of plant diseases on a wide range of crops. (See J. R. Godwin, V. M. Anthony, J. M. Clough and C. R. A. Godfrey "ICIA5504: A Novel, Broad Spectrum, Systemic β-Methoxyacrylate Fungicide" in Brighton Crop Protection Conference - Pests and Diseases - 1992, 1, 435-442.) European Patent Application EP-A-398,692 discloses (E)-2-(methoxyimino)-N-methyl-2-(2-phenoxyphenyl)acetamide also known as SSF 126. This fungicide is under consideration for the control of plant disease, especially of rice. EP-A-68813 discloses certain triazole compounds as fungicides (e.g., flusilazole). EP-A-40345 discloses certain triazole compounds as fungicides (e.g., tebuconazole).

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SUMMARY OF THE INVENTION

This invention is directed to fungicidal combinations (e.g., mixtures) comprising (1) at least one compound selected from the quinazolinones of Formula I (including all geometric and stereoisomers), *N*-oxides, and agriculturally suitable salts thereof,

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wherein

D is O or S:

 R^{\dagger} is C_1 - C_{10} alkyl; C_3 - C_5 cycloalkyl; C_3 - C_{10} alkenyl; C_3 - C_{10} alkynyl;

 C_1 - C_{10} haloalkyl; C_3 - C_{10} haloalkenyl; C_3 - C_{10} haloalkynyl;

C₂-C₁₀ alkoxyalkyl; C₂-C₁₀ alkylthioalkyl; C₂-C₁₀ alkylsulfinylalkyl;

C₂-C₁₀ alkylsulfonylalkyl; C₄-C₁₀ cycloalkylalkyl;

C₄-C₁₀ alkenyloxyalkyl; C₄-C₁₀ alkynyloxyalkyl; C₄-C₁₀

(cycloalkyl)oxyalkyl; C₄-C₁₀ alkenylthioalkyl; C₄-C₁₀ alkynylthioalkyl;

C₄-C₁₀ (cycloalkyl)thioalkyl; C₂-C₁₀ haloalkoxyalkyl;

C₄-C₁₀ haloalkenyloxyalkyl; C₄-C₁₀ haloalkynyloxyalkyl;

C₄-C₁₀ alkoxyalkenyl; C₄-C₁₀ alkoxyalkynyl; C₄-C₁₀ alkylthioalkenyl;

 C_4 - C_{10} alkylthioalkynyl; C_4 - C_{10} trialkylsilylalkyl; C_1 - C_{10} alkyl substituted with NR⁵R⁶, nitro, cyano or phenyl optionally substituted with R⁸, R⁹ and

R¹⁰; C₁-C₁₀ alkoxy; C₁-C₁₀ haloalkoxy; C₁-C₁₀ alkylthio;

C₁-C₁₀ haloalkylthio; or pyridinyl, furanyl, thienyl, naphthalenyl,

benzofuranyl, benzothienyl or quinolinyl each optionally substituted with

R8, R9 and R10;

 $R^2 \text{ is } C_1\text{-}C_{10} \text{ alkyl}; C_3\text{-}C_7 \text{ cycloalkyl}; C_3\text{-}C_{10} \text{ alkenyl}; C_3\text{-}C_{10} \text{ alkynyl};\\$

 C_1 - C_{10} haloalkyl; C_3 - C_{10} haloalkenyl; C_3 - C_{10} haloalkynyl;

C₂-C₁₀ alkoxyalkyl; C₂-C₁₀ alkylthioalkyl; C₂-C₁₀ alkylsulfinylalkyl;

C₂-C₁₀ alkylsulfonylalkyl; C₄-C₁₀ cycloalkylalkyl;

 C_4 - C_{10} alkenyloxyalkyl; C_4 - C_{10} alkynyloxyalkyl; C_4 - C_{10}

(cycloalkyl)oxyalkyl; C₄-C₁₀ alkenylthioalkyl; C₄-C₁₀ alkynylthioalkyl;

C₄-C₁₀ (cycloalkyl)thioalkyl; C₂-C₁₀ haloalkoxyalkyl;

C₄-C₁₀ haloalkenyloxyalkyl; C₄-C₁₀ haloalkynyloxyalkyl;

 C_4 - C_{10} alkoxyalkenyl; C_4 - C_{10} alkoxyalkynyl; C_4 - C_{10} alkylthioalkenyl;

C₄-C₁₀ alkylthioalkynyl; C₄-C₁₀ trialkylsilylalkyl; C₂-C₁₀ cyanoalkyl;

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 C_1 - C_{10} nitroalkyl; C_1 - C_{10} alkyl substituted with CO_2R^5 , NR^5R^6 , or phenyl optionally substituted with R^7 , R^9 and R^{10} ; phenyl optionally substituted with R^7 , R^9 and R^{10} ; -N= CR^5R^5 ; or -NR $^5R^6$; or

 R^1 and R^2 are taken together to form -CH₂(CH₂)_qCH₂-;

5 q is 0, 1, 2, 3 or 4;

R³ is halogen, C₁-C₈ alkyl, C₃-C₈ cycloalkyl, C₂-C₈ alkenyl, C₂-C₈ alkynyl, C₁-C₈ haloalkyl, C₃-C₈ haloalkenyl, C₃-C₈ haloalkynyl, C₁-C₈ alkoxy, C₁-C₈ haloalkoxy, C₃-C₈ alkenyloxy, C₃-C₈ alkynyloxy, C₁-C₈ alkylthio, C₃-C₈ alkenylthio, C₁-C₈ alkylsulfinyl, C₁-C₈ alkylsulfinyl, C₁-C₈ alkylsulfinyl, C₂-C₈ alkoxyalkyl, C₂-C₈ alkylsulfinylalkyl, C₂-C₈ alkylsulfonylalkyl, C₄-C₈ cycloalkylalkyl, C₃-C₈ trialkylsilyl, NR⁵R⁶, C₅-C₈ trialkylsilylalkynyl or phenyl optionally substituted with at least one R⁷;

 R^4 is hydrogen, halogen, C_1 - C_4 alkyl, C_1 - C_4 haloalkyl, C_1 - C_4 alkoxy or C_1 - C_4 haloalkoxy;

each R^5 is independently hydrogen, C_1 - C_4 alkyl or phenyl optionally substituted with at least one R^7 ;

each R^6 is independently hydrogen, C_1 - C_8 alkyl or phenyl optionally substituted with at least one R^7 ; or

when R⁵ and R⁶ are attached to the same nitrogen atom, R⁵ and R⁶ can be taken together to form -CH₂CH₂CH₂CH₂-, -CH₂(CH₂)₃CH₂-, -CH₂CH₂OCH₂CH₂-, -CH₂CH(Me)CH₂- or -CH₂CH(Me)OCH(Me)CH₂-;

each R⁷ is independently halogen, C₁-C₄ alkyl, C₁-C₄ alkoxy, C₁-C₄ haloalkyl, nitro or cyano;

 R^8 is C_1 - C_6 alkyl, C_1 - C_6 alkoxy, C_1 - C_6 haloalkyl, halogen, C_2 - C_8 alkynyl, C_1 - C_6 alkylthio, phenyl or phenoxy each optionally substituted with at least one R^7 , cyano, nitro, C_1 - C_6 haloalkoxy, C_1 - C_6 haloalkylthio, C_2 - C_6 alkenyl, C_2 - C_6 haloalkenyl, acetyl, CO_2 Me or $N(C_1$ - C_2 alkyl)₂;

each R⁹ is independently methyl, ethyl, methoxy, methylthio, halogen or trifluoromethyl; and

each R¹⁰ is independently halogen;

and (2) at least one compound selected from (a) compounds acting at the bc_1 complex of the fungal mitochondrial respiratory electron transfer site and (b) compounds that control fungal disease by inhibiting the sterol biosynthesis pathway. This invention provides agricultural compositions containing these combinations and the use of the combinations as fungicides. Advantageous compositions include those where component (1) and component (2) are present in a fungicidally effective amount and the

mole ratio of component (1) to component (2) is from about 30:1 to 1:30. Advantageous methods include those where component (1) and component (2) are added in amounts sufficient to provide a fungicidal effectiveness greater than the sum of the fungicidal effectivenesses provided by those amounts of said components taken independently.

DETAILS OF THE INVENTION

Combinations of fungicides are used in accordance with this invention to facilitate disease control and to retard resistance development. Suitable compositions and methods are provided.

For example, this invention also provides methods for controlling plant diseases caused by fungal plant pathogens comprising applying to the plant or portion thereof to be protected, or to the plant seed or seedling to be protected one of the following:

A) an effective amount of a fungicidal composition comprising component (1), component (2), and at least one of a surfactant, a solid diluent or a liquid diluent;

B) (i) an effective amount of a first composition comprising component (1), and at least one of a surfactant, solid or liquid diluent; and (ii) an effective amount of a second composition comprising component (2), and at least one of a surfactant, a solid diluent or a liquid diluent; said first and second compositions applied sequentially in any order; or

C) an effective amount of a physical mixture of the first and second compositions as defined in B above.

The mole ratio of the compound(s) of component (1) to the compound(s) of component (2) applied is normally from about 30:1 to 1:30, and the compound(s) of component (1) and the compound(s) of component (2) are normally applied in amounts effective to provide control of the fungal disease which is greater than the additive control of that fungal disease provided by the compound(s) of component (1) and the compound(s) of component (2) individually.

Preferred compositions for reasons of ease of synthesis or greater fungicidal activity are:

Preferred 1. A fungicidal composition comprising a fungicidally effective amount of (1) at least one quinazolinone of Formula I (including all geometric and stereoisomers), *N*-oxides, and agriculturally suitable salts thereof, wherein

$$R^1$$
 is C_1 - C_6 alkyl;

$$R^2$$
 is C_1 - C_6 alkyl;

R³ is halogen and fixed at position 6 of the quinazolinone; and R⁴ is hydrogen or halogen and fixed at position 8 of the quinazolinone; and (2) at least one compound selected from

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(a) compounds of Formula II (including all geometric and stereoisomers), *N*-oxides, and agriculturally suitable salts thereof,

H

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wherein

Q is

$$X \xrightarrow{G} W$$

or

 $A = N$
 R^{12}
 $Q-1$
 $Q-2$

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E is selected from:

- i) 1,2-phenylene optionally substituted with R^{13} or both R^{13} and R^{14} ;
- ii) a naphthalene ring, provided that when G and Y are attached to the same ring, then G and Y are attached to adjacent ring members, the naphthalene ring optionally substituted with R¹³ or both R¹³ and R¹⁴; and

iii) a ring system selected from 5 to 12-membered monocyclic and fused bicyclic aromatic heterocyclic ring systems, each heterocyclic ring system containing 1 to 6 heteroatoms independently selected from the group nitrogen, oxygen, and sulfur, provided that each heterocyclic ring system contains no more than 4 nitrogens, no more than 2 oxygens, and no more than 2 sulfurs, each fused bicyclic ring system optionally containing one nonaromatic ring that optionally includes one or two J¹ as ring members and optionally includes one or two ring members independently selected from C(=O) and S(O)₂, provided that G is attached to an aromatic ring, and when G and Y are attached to the same ring, then G and Y are attached to adjacent ring members, each aromatic

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heterocyclic ring system optionally substituted with R¹³ or both R¹³ and R¹⁴; A is O, S, N, NR¹⁵ or CR²⁴; G is C or N; provided that when G is C, then A is O, S or NR¹⁵ and the floating double bond is attached to G; and when G is N, then A is N 5 or CR²⁴ and the floating double bond is attached to A; W is O, S, NH, $N(C_1-C_6 \text{ alkyl})$ or $NO(C_1-C_6 \text{ alkyl})$; W¹ is O or NH; X is H, OR^{11} , $S(O)_mR^{11}$, halogen, C_1 - C_6 alkyl, C_1 - C_6 haloalkyl, C₃-C₆ cycloalkyl, cyano, NH₂, NHR¹¹, N(C₁-C₆ alkyl)R¹¹, 10 $NH(C_1-C_6 \text{ alkoxy}) \text{ or } N(C_1-C_6 \text{ alkoxy})R^{11};$ X¹ is CH or N; R¹¹ is C₁-C₆ alkyl, C₁-C₆ haloalkyl, C₂-C₆ alkenyl, C₂-C₆ haloalkenyl, C2-C6 alkynyl, C2-C6 haloalkynyl, C3-C6 cycloalkyl, C₂-C₄ alkylcarbonyl or C₂-C₄ alkoxycarbonyl; 15 R¹² is H, C₁-C₆ alkyl, C₁-C₆ haloalkyl, C₂-C₆ alkenyl, C₂-C₆ haloalkenyl, C₂-C₆ alkynyl, C₂-C₆ haloalkynyl, C₃-C₆ cycloalkyl, C₂-C₄ alkylcarbonyl, C₂-C₄ alkoxycarbonyl, hydroxy, C₁-C₂ alkoxy or acetyloxy; R¹³ and R¹⁴ are each independently halogen; cyano; nitro; hydroxy; 20 C₁-C₆ alkyl; C₁-C₆ haloalkyl; C₂-C₆ alkenyl; C₂-C₆ haloalkenyl; C_2 - C_6 alkynyl; C_2 - C_6 haloalkynyl; C_1 - C_6 alkoxy; C₁-C₆ haloalkoxy; C₂-C₆ alkenyloxy; C₂-C₆ alkynyloxy; C₁-C₆ alkylthio; C₁-C₆ alkylsulfinyl; C₁-C₆ alkylsulfonyl; formyl; C₂-C₆ alkylcarbonyl; C₂-C₆ alkoxycarbonyl; NH₂C(O); 25 $(C_1-C_4 \text{ alkyl})NHC(O); (C_1-C_4 \text{ alkyl})_2NC(O); Si(R^{35})_3; Ge(R^{35})_3;$ (R³⁵)₃Si-C≡C-; or phenyl, phenylethynyl, benzoyl, or phenylsulfonyl each substituted with R18 and optionally substituted with one or more R^{20} ; or when E is 1,2-phenylene and R¹³ and R¹⁴ are attached to adjacent atoms, 30 R¹³ and R¹⁴ can be taken together as C₃-C₅ alkylene, C₃-C₅ haloalkylene, C₃-C₅ alkenylene or C₃-C₅ haloalkenylene each optionally substituted with 1-2 C₁-C₃ alkyl; R^{15} is H, C_1 - C_6 alkyl, C_1 - C_6 haloalkyl, C_2 - C_6 alkenyl, C2-C6 haloalkenyl, C2-C6 alkynyl, C2-C6 haloalkynyl, 35 C₃-C₆ cycloalkyl, C₂-C₄ alkylcarbonyl or C₂-C₄ alkoxycarbonyl; Y is -O-, -S(O)_n-, -NR²⁵-, -C(=O)-, -CH(OR²⁵)-, -CHR¹⁶-, -CHR¹⁶CHR¹⁶-, -CR¹⁶=CR¹⁶-, -C=C-, -CHR²⁵O-, -OCHR²⁵-,

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-CHR<sup>25</sup>S(O)<sub>n</sub>-, -S(O)<sub>n</sub>CHR<sup>25</sup>-, -CHR<sup>25</sup>O-N=C(R<sup>17</sup>)-,
                                   -(R^{17})C=N-OCH(R^{25})-, -C(R^{17})=N-O-, -O-N=C(R^{17})-,
                                   -CHR^{25}OC(=O)N(R^{25})-, -CHR^{25}OC(=S)N(R^{25})-,
                                   -CHR<sup>25</sup>OC(=O)O-, -CHR<sup>25</sup>OC(=S)O-, -CHR<sup>25</sup>OC(=O)S-,
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                                   -CHR^{25}OC(=S)S_{-}, -CHR^{25}SC(=O)N(R^{25})_{-},
                                   -CHR^{25}SC(=S)N(R^{25})-, -CHR^{25}SC(=O)O-, -CHR^{25}SC(=S)O-.
                                   -CHR<sup>25</sup>SC(=O)S-, -CHR<sup>25</sup>SC(=S)S-, -CHR<sup>25</sup>SC(=NR<sup>25</sup>)S-,
                                   -CHR^{25}N(R^{25})C(=O)N(R^{25})-, -CHR^{25}O-N(R^{25})C(=O)N(R^{25})-,
                                   -CHR^{25}O-N(R^{25})C(=S)N(R^{25})-, -CHR^{25}O-N=C(R^{17})NR^{25}-,
                                   -CHR^{25}O-N=C(R^{17})OCH_{2-}, -CHR^{25}O-N=C(R^{17})-N=N-
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                                   -CHR^{25}O-N=C(R^{17})-C(=O)-
                                   -CHR^{25}O-N=C(R^{17})-C(=N-A^2-Z^1)-A^1
                                   -CHR^{25}O-N=C(R^{17})-C(R^{17})=N-A^2-A^3
                                   -CHR^{25}O-N=C(-C(R^{17})=N-A^2-Z^1)-, -CHR^{25}O-N=C(R^{17})-CH_2O-,
                                   -CHR<sup>25</sup>O-N=C(R<sup>17</sup>)-CH<sub>2</sub>S-, -O-CH<sub>2</sub>CH<sub>2</sub>O-N=C(R<sup>17</sup>)-,
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                                   -CHR^{25}O-C(R^{25})=C(R^{17})-, -CHR^{25}O-C(R^{17})=N-,
                                   -CHR^{25}S-C(R^{17})=N-, -C(R^{17})=N-NR^{25}-, -CH=N-N=C(R^{17})-,
                                   -CHR^{25}N(R^{25})-N=C(R^{17})-, -CHR^{25}N(COCH_3)-N=C(R^{17})-,
                                   -OC(=S)NR^{25}C(=O)-, -CHR^{16}-C(=W^2)-A^1-,
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                                   -CHR^{16}CHR^{16}-C(=W^2)-A^1-, -CR^{16}=CR^{16}-C(=W^2)-A^1-,
                                   -C = C - C(=W^2) - A^1 -, -N = CR^{16} - C(=W^2) - A^1 - or a direct bond; and
                                   the directionality of the Y linkage is defined such that the moiety
                                   depicted on the left side of the linkage is bonded to E and the
                                   moiety on the right side of the linkage is bonded to Z;
25
                           Z^1 is H or -A<sup>3</sup>-Z;
                            W<sup>2</sup> is O or S:
                            A<sup>1</sup> is O, S, NR<sup>25</sup> or a direct bond:
                            A<sup>2</sup> is O, NR<sup>25</sup> or a direct bond;
                            A^3 is -C(=O)-, -S(O)_2- or a direct bond;
                           each R<sup>16</sup> is independently H, 1-2 CH<sub>3</sub>, C<sub>2</sub>-C<sub>3</sub> alkyl, C<sub>1</sub>-C<sub>3</sub> alkoxy,
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                                   C_3-C_6 cycloalkyl, formylamino, C_2-C_4 alkylcarbonylamino,
                                   C<sub>2</sub>-C<sub>4</sub> alkoxycarbonylamino, NH<sub>2</sub>C(O)NH,
                                   (C<sub>1</sub>-C<sub>3</sub> alkyl)NHC(O)NH, (C<sub>1</sub>-C<sub>3</sub> alkyl)<sub>2</sub>NC(O)NH,
                                   N(C<sub>1</sub>-C<sub>3</sub> alkyl)<sub>2</sub>, piperidinyl, morpholinyl, 1-2 halogen, cyano or
35
                                   nitro;
                            each R<sup>17</sup> is independently H, C<sub>1</sub>-C<sub>6</sub> alkyl, C<sub>1</sub>-C<sub>6</sub> haloalkyl,
                                    C_1-C_6 alkoxy, C_1-C_6 haloalkoxy, C_1-C_6 alkylthio,
                                    C<sub>1</sub>-C<sub>6</sub> alkylsulfinyl, C<sub>1</sub>-C<sub>6</sub> alkylsulfonyl, C<sub>1</sub>-C<sub>6</sub> haloalkylthio,
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C₁-C₆ haloalkylsulfinyl, C₁-C₆ haloalkylsulfonyl, C₂-C₆ alkenyl, C2-C6 haloalkenyl, C2-C6 alkynyl, C2-C6 haloalkynyl, C₃-C₆ cycloalkyl, C₂-C₄ alkylcarbonyl, C₂-C₄ alkoxycarbonyl, halogen, cyano, nitro, hydroxy, amino, NH(C₁-C₆ alkyl), $N(C_1-C_6 \text{ alkyl})_2$ or morpholinyl; each Z is independently selected from: i) C₁-C₁₀ alkyl, C₂-C₁₀ alkenyl and C₂-C₁₀ alkynyl each substituted with R¹⁹ and optionally substituted with one or more R²⁰: ii) C3-C8 cycloalkyl, C3-C8 cycloalkenyl and phenyl each substituted with R¹⁹ and optionally substituted with one or more R^{20} ; iii) a ring system selected from 3 to 14-membered monocyclic, fused bicyclic and fused tricyclic nonaromatic heterocyclic ring systems and 5 to 14-membered monocyclic, fused bicyclic and fused tricyclic aromatic heterocyclic ring systems, each heterocyclic ring system containing 1 to 6 heteroatoms independently selected from the group nitrogen, oxygen, and sulfur, provided that each heterocyclic ring system contains no more than 4 nitrogens, no more than 2 oxygens, and no more than 2 sulfurs, each nonaromatic or aromatic heterocyclic ring system substituted with R^{19} and optionally substituted with one or more R^{20} ; iv) a multicyclic ring system selected from 8 to 14-membered fused-bicyclic and fused-tricyclic ring systems which are an aromatic carbocyclic ring system, a nonaromatic carbocyclic ring system, or a ring system containing one or two nonaromatic rings that each include one or two J1 as ring members and one or two ring members independently selected from C(=O) and S(O)2, and any remaining rings as aromatic carbocyclic rings, each multicyclic ring system substituted with R19 and optionally substituted with one or more R20; and v) adamantyl substituted with R¹⁹ and optionally substituted with one or more R²⁰;

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each J1 is independently selected from the group -CHR²³-, -NR²³-, -O-

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 R^{18} is H, 1-2 halogen, C_1 - C_6 alkyl, C_1 - C_6 haloalkyl, C_1 - C_6 alkoxy, C₁-C₆ haloalkoxy, C₂-C₆ alkenyl, C₂-C₆ haloalkenyl, C2-C6 alkynyl, C1-C6 alkylthio, C1-C6 haloalkylthio,

and $-S(O)_{p}$ -;

	C_1 - C_6 arkyrsuninyr, C_1 - C_6 arkyrsunionyr, C_3 - C_6 cycloarkyr,
	C_3 - C_6 alkenyloxy, $CO_2(C_1$ - C_6 alkyl), $NH(C_1$ - C_6 alkyl),
	$N(C_1-C_6 \text{ alkyl})_2$, cyano, nitro, $SiR^{29}R^{30}R^{31}$ or $GeR^{29}R^{30}R^{31}$;
	R^{19} is H; 1-2 halogen; C_1 - C_6 alkyl; C_1 - C_6 haloalkyl; C_1 - C_6 alkoxy;
5	C ₁ -C ₆ haloalkoxy; C ₂ -C ₆ alkenyl; C ₂ -C ₆ haloalkenyl;
	C ₂ -C ₆ alkynyl; C ₁ -C ₆ alkylthio; C ₁ -C ₆ haloalkylthio;
	C ₁ -C ₆ alkylsulfinyl; C ₁ -C ₆ alkylsulfonyl; C ₃ -C ₆ cycloalkyl;
	C_3 - C_6 alkenyloxy; $CO_2(C_1$ - C_6 alkyl); $NH(C_1$ - C_6 alkyl);
	$N(C_1-C_6 \text{ alkyl})_2$; $-C(R^{28})=NOR^{27}$; cyano; nitro; SF_5 ;
10	$SiR^{32}R^{33}R^{34}$; or $GeR^{32}R^{33}R^{34}$; or R^{19} is phenyl, benzyl, benzyl,
	phenoxy, pyridinyl, pyridinyloxy, thienyl, thienyloxy, furanyl,
	pyrimidinyl or pyrimidinyloxy each optionally substituted with R ²¹
	or both R^{21} and R^{22} ;
	each R ²⁰ is independently halogen, C ₁ -C ₄ alkyl, C ₁ -C ₄ haloalkyl,
15	C ₁ -C ₄ alkoxy, nitro or cyano; or
	when R ¹⁹ and an R ²⁰ are attached to adjacent atoms on Z, R ¹⁹ and said
	adjacently attached R^{20} can be taken together as -OCH ₂ O- or
	-OCH ₂ CH ₂ O-; each CH ₂ group of said taken together R ¹⁹ and R ²⁰
	optionally substituted with 1-2 halogen; or
20	when Y and an R ²⁰ are attached to adjacent atoms on Z and Y is
	-CHR ²⁵ O-N=C(R ¹⁷)-, -O-N=C(R ¹⁷)-, -O-CH ₂ CH ₂ O-N=C(R ¹⁷)-,
	-CHR ²⁵ O-C(R ²⁵)=C(R ¹⁷)-, -CH=N-N=C(R ¹⁷)-,
	-CHR ²⁵ N(R ²⁵)-N=C(R ¹⁷)- or -CHR ²⁵ N(COCH ₃)-N=C(R ¹⁷)-, R ¹⁷
	and said adjacently attached R ²⁰ can be taken together as
25	- $(CH_2)_r$ -J- such that J is attached to Z;
	J is -CH ₂ -, -CH ₂ CH ₂ -, -OCH ₂ -, -CH ₂ O-, -SCH ₂ -, -CH ₂ S-, -N(\mathbb{R}^{26})CH ₂ -
	or -CH ₂ N(R ²⁶)-; each CH ₂ group of said J optionally substituted
•	with 1 to 2 CH ₃ ;
	R^{21} and R^{22} are each independently 1-2 halogen; C_1 - C_4 alkyl;
30	C ₁ -C ₄ haloalkyl; C ₂ -C ₆ alkenyl; C ₂ -C ₆ haloalkenyl; C ₂ -C ₆
	alkynyl; C ₂ -C ₆ haloalkynyl; C ₂ -C ₆ alkoxyalkyl; C ₂ -C ₆
	alkylthioalkyl; C ₃ -C ₆ alkoxyalkynyl;
	C ₇ -C ₁₀ tetrahydropyranyloxyalkynyl; benzyloxymethyl; C ₁ -C ₄
	alkoxy; C ₁ -C ₄ haloalkoxy; C ₃ -C ₆ alkenyloxy; C ₃ -C ₆
35	haloalkenyloxy; C ₃ -C ₆ alkynyloxy; C ₃ -C ₆ haloalkynyloxy; C ₂ -C ₆
	alkoxyalkoxy; C5-C9 trialkylsilylalkoxyalkoxy;
	C ₂ -C ₆ alkylthioalkoxy; C ₁ -C ₄ alkylthio; C ₁ -C ₄ haloalkylthio;
	C ₁ -C ₄ alkylsulfinyl; C ₁ -C ₄ haloalkylsulfinyl; C ₁ -C ₄ alkylsulfonyl;

	C_1 - C_4 haloalkylsulfonyl; C_3 - C_6 alkenylthio;
	C ₃ -C ₆ haloalkenylthio; C ₂ -C ₆ alkylthioalkylthio; nitro; cyano;
•	thiocyanato; hydroxy; $N(R^{36})_2$; SF_5 ; $Si(R^{35})_3$; $Ge(R^{35})_3$;
	$(R^{35})_3$ Si-C=C-; $OSi(R^{35})_3$; $OGe(R^{35})_3$; $C(=O)R^{36}$; $C(=S)R^{36}$;
5	$C(=O)OR^{36}$; $C(=S)OR^{36}$; $C(=O)SR^{36}$; $C(=S)SR^{36}$; $C(=O)N(R^{36})_2$;
	$C(=S)N(R^{36})_2$; $OC(=O)R^{36}$; $OC(=S)R^{36}$; $SC(=O)R^{36}$; $SC(=S)R^{36}$;
	$N(R^{36})C(=O)R^{36}; N(R^{36})C(=S)R^{36}; OC(=O)OR^{37}; OC(=O)SR^{37};$
	$OC(=O)N(R^{36})_2$; $SC(=O)OR^{37}$; $SC(=O)SR^{37}$; $S(O)_2OR^{36}$;
	$S(O)_2N(R^{36})_2$; $OS(O)_2R^{37}$; $N(R^{36})S(O)_2R^{37}$; or phenyl, phenoxy,
10	benzyl, benzyloxy, phenylsulfonyl, phenylethynyl or
	pyridinylethynyl, each optionally substituted with halogen,
	C_1 - C_4 alkyl, C_1 - C_4 haloalkyl, C_1 - C_4 alkoxy, C_1 - C_4 haloalkoxy,
	nitro or cyano;
	each R ²³ is independently H; C ₁ -C ₆ alkyl; C ₁ -C ₆ haloalkyl; or phenyl
15	optionally substituted with halogen, C_1 - C_4 alkyl, C_1 - C_4 haloalkyl,
	C ₁ -C ₄ alkoxy, C ₁ -C ₄ haloalkoxy, nitro or cyano;
	R^{24} is H, halogen, C_1 - C_6 alkyl, C_1 - C_6 haloalkyl, C_2 - C_6 alkenyl,
	C ₂ -C ₆ haloalkenyl, C ₂ -C ₆ alkynyl, C ₂ -C ₆ haloalkynyl or
	C ₃ -C ₆ cycloalkyl;
20	each R ²⁵ is independently H; C ₁ -C ₃ alkyl; C ₃ -C ₆ cycloalkyl; or phenyl or
	benzyl, each optionally substituted on the phenyl ring with halogen,
	C_1 - C_4 alkyl, C_1 - C_4 haloalkyl, C_1 - C_4 alkoxy, C_1 - C_4 haloalkoxy,
	nitro or cyano; or
25	when Y is $-CHR^{25}N(R^{25})C(=O)N(R^{25})$ -, the two R^{25} attached to
25	nitrogen atoms on said group can be taken together as - $(CH_2)_S$ -; or
	when Y is -CHR ²⁵ O-N=C(R ¹⁷)NR ²⁵ -, R ¹⁷ and the adjacently attached
	R^{25} can be taken together as $-CH_2-(CH_2)_S$ -, $-O-(CH_2)_S$ -,
	-S- $(CH_2)_{S}$ - or -N $(C_1-C_3 \text{ alkyl})$ - $(CH_2)_{S}$ -; with the directionality of said linkage defined such that the moiety depicted on the left side
30	of the linkage is bonded to the carbon and the moiety on the right
30	side of the linkage is bonded to the nitrogen;
	R^{26} , R^{27} and R^{28} are each independently H; C_1 - C_3 alkyl;
	C ₃ -C ₆ cycloalkyl; or phenyl optionally substituted with halogen,
	C_1 - C_4 alkyl, C_1 - C_4 haloalkyl, C_1 - C_4 alkoxy, C_1 - C_4 haloalkoxy,
35	nitro or cyano;
	R^{29} , R^{30} , R^{31} , R^{32} , R^{33} and R^{34} are each independently C_1 - C_6 alkyl,
	C_2 - C_6 alkenyl, C_1 - C_4 alkoxy or phenyl;

	each R^{35} is independently C_1 - C_4 alkyl, C_1 - C_4 haloalkyl, C_2 - C_4 alkenyl,
	C ₁ -C ₄ alkoxy or phenyl;
	each R ³⁶ is independently H; C ₁ -C ₆ alkyl; C ₁ -C ₆ haloalkyl;
	C_2 - C_6 alkenyl; C_2 - C_6 haloalkenyl; C_2 - C_6 alkynyl;
5	C_2 - C_6 haloalkynyl; C_3 - C_6 cycloalkyl; or phenyl or benzyl, each
	optionally substituted on the phenyl ring with halogen, C ₁ -C ₄ alkyl,
	C_1 - C_4 haloalkyl, C_1 - C_4 alkoxy, C_1 - C_4 haloalkoxy, nitro or cyano;
	each R^{37} is independently C_1 - C_6 alkyl; C_1 - C_6 haloalkyl; C_2 - C_6 alkenyl;
	C ₂ -C ₆ haloalkenyl; C ₂ -C ₆ alkynyl; C ₂ -C ₆ haloalkynyl;
10	C ₃ -C ₆ cycloalkyl; or phenyl or benzyl, each optionally substituted
	on the phenyl ring with halogen, C_1 - C_4 alkyl, C_1 - C_4 haloalkyl,
	C ₁ -C ₄ alkoxy, C ₁ -C ₄ haloalkoxy, nitro or cyano;
	m, n and p are each independently 0, 1 or 2;
1.5	r is 0 or 1; and
15	s is 2 or 3;
	and (b) bromuconazole, cyproconazole, difenoconazole, diniconazole,
	epoxiconazole, fenarimol, fenbuconazole, fenpropidin,
	fenpropimorph, fluquinconazole, flusilazole, flutriafol,
20	hexaconazole, ipconazole, metconazole, penconazole,
20	propiconazole, tebuconazole, tetraconazole, triadimefon,
	triadimenol, tridemorph, triticonazole and uniconazole.
	Preferred 2. The fungicidal composition of Preferred 1 comprising a fungicidally effective amount of (1) at least one compound selected from the group
	consisting of
25	6-bromo-3-propyl-2-propyloxy-4(3 <i>H</i>)-quinazolinone,
4 5	6,8-diiodo-3-propyl-2-propyloxy-4(3 <i>H</i>)-quinazolinone and
	6-iodo-3-propyl-2-propyloxy-4(3H)-quinazolinone;
	and (2) at least one compound selected from the group consisting of
	2,4-dihydro-5-methoxy-2-methyl-4-[2-[[[[1-[3-
30	(trifluoromethyl)phenyl]ethylidenc]amino]oxy]methyl]
	phenyl]- $3H$ -1,2,4-triazol-3-one, kresoxim-methyl,
	azoxystrobin, (E)-2-(methoxyimino)-N-methyl-2-(2-
	phenoxyphenyl)acetamide, flusilazole, epoxiconazole,
	fenpropimorph, propiconazole and tebuconazole.
35	Preferred 3. The fungicidal composition of Preferred 2 comprising a fungicidally
	effective amount of (1) 6-iodo-3-propyl-2-propyloxy-4(3H)-quinazolinone
	(sometimes referred to hereafter as the Formula Ia compound) and (2) 2,4-
	dihydro-5-methoxy-2-methyl-4-[2-[[[[1-[3-

(trifluoromethyl)phenyl]ethylidene]amino]oxy]methyl]phenyl]-3*H*-1,2,4-triazol-3-one (sometimes referred to hereafter as the Formula IIa compound).

Preferred 4. The fungicidal composition of Preferred 2 comprising a fungicidally effective amount of (1) 6-iodo-3-propyl-2-propyloxy-4(3*H*)-quinazolinone and (2) flusilazole.

Preferred 5. The fungicidal composition of Preferred 2 comprising a fungicidally effective amount of (1) 6-iodo-3-propyl-2-propyloxy-4(3*H*)-quinazolinone and (2) tebuconazole.

Preferred 6. The fungicidal composition of Preferred 2 comprising a fungicidally effective amount of (1) 6-iodo-3-propyl-2-propyloxy-4(3*H*)-quinazolinone, and (2) both 2,4-dihydro-5-methoxy-2-methyl-4-[2-[[[1-[3-(trifluoromethyl)phenyl]ethylidene]amino]oxy]methyl]phenyl]-3*H*-1,2,4-triazol-3-one and flusilazole.

In the above recitations, the term "alkyl", used either alone or in compound words such as "alkylthio" or "haloalkyl" includes straight-chain or branched alkyl, such as, methyl, ethyl, n-propyl, i-propyl, or the different butyl, pentyl, hexyl, heptyl, octyl, nonyl and decyl isomers. The term "1-2 CH₃" indicates that the substituent can be methyl (i.e., Me) or, when there is a hydrogen attached to the same atom, the substituent and said hydrogen can both be methyl. "Alkenyl" includes straight-chain or branched alkenes such as ethenyl, 1-propenyl, 2-propenyl, and the different butenyl, pentenyl, hexenyl, heptenyl, octenyl, nonenyl and decenyl isomers. "Alkenyl" also includes polyenes such as 1,2-propadienyl and 2,4-hexadienyl. "Alkynyl" includes straight-chain or branched alkynes such as ethynyl, 1-propynyl, 2-propynyl and the different butynyl, pentynyl, hexynyl, heptynyl, octynyl, nonynyl, and decynyl isomers. "Alkynyl" can also include moieties comprised of multiple triple bonds such as 2,5-hexadiynyl.

"Alkylene" denotes a straight-chain or branched alkanediyl. Examples of "alkylene" include CH₂CH₂CH₂, CH₂CH(CH₃) and the different butylene and pentylene isomers. "Haloalkylene" denotes a halogen substituted alkylene. Examples of "haloalkylene" include CH₂CH(CF₃), CH₂CF₂CH₂ and CH₂CH(CCl₃). "Alkenylene" denotes a straight-chain or branched alkenediyl containing one olefinic bond. Examples of "alkenylene" include CH₂CH=CH and CH=C(CH₃) and the different butenylene and pentenylene isomers. "Haloalkenylene" denotes a halogen substituted alkenylene. Examples of "haloalkenylene" include CH₂CCl=CCl and CH=C(CF₃).

"Alkenyloxyalkyl" denotes alkenyl substitution on oxygen which in turn is substituted on alkyl. Examples "alkenyloxyalkyl" include CH₂=CHCH₂OCH₂ and CH₃CH=CHCH₂OCH₂CH₂. "Alkynyloxyalkyl" denotes alkynyl substitution on

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oxygen which in turn is substituted on alkyl. Examples of "alkynyloxyalkyl" include CH≡CCH₂OCH₂ and CH₃C≡CCH₂OCH₂CH₂.

"Alkoxy" includes, for example, methoxy, ethoxy, propyloxy, 1-methylethoxy and the different butoxy, pentyloxy, hexyloxy, heptyloxy, octyloxy, nonyloxy, and 5 decyloxy isomers. "Alkoxyalkyl" denotes alkoxy substitution on alkyl. Examples of "alkoxyalkyl" include CH₃OCH₂, CH₃OCH₂CH₂, CH₃CH₂OCH₂, CH₃CH₂CH₂OCH₂ and CH₃CH₂OCH₂CH₂. "Alkoxyalkoxy" denotes alkoxy substitution on alkoxy. Examples of "alkoxyalkoxy" include CH₃OCH₂O, (CH₃)₃COCH₂O and CH₃OCH₂CH₂O. "Alkenyloxy" includes straight-chain or 10 branched alkenyloxy moieties. Examples of "alkenyloxy" include H₂C=CHCH₂O, (CH₃)₂C=CHCH₂O, (CH₃)CH=CHCH₂O, (CH₃)CH=C(CH₃)CH₂O and CH₂=CHCH₂CH₂O. "Alkynyloxy" includes straight-chain or branched alkynyloxy moieties. Examples of "alkynyloxy" include HC=CCH2O, CH3C=CCH2O and CH₃C≡CCH₂CH₂O. "Alkoxyalkenyl" denotes alkoxy substitution of alkenyl. "Alkoxyalkenyl" includes straight-chain or branched alkoxyalkenyl moieties. Examples 15 of "alkoxyalkenyl" include (CH₃)₂CHOCH=CH and CH₃OCH₂CH=CH. "Alkoxyalkynyl" denotes alkoxy substitution of alkynyl. "Alkoxyalkynyl" includes straight-chain or branched alkoxyalkynyl moieties. Examples of "alkoxyalkynyl"

"Alkylthio" includes branched or straight-chain alkylthio moieties such as methylthio, ethylthio, and the different propylthio, butylthio, pentylthio, hexylthio, heptylthio, octylthio, nonylthio and decylthio isomers. "Alkylthioalkyl" denotes alkylthio substitution on alkyl. Examples of "alkylthioalkyl" include CH₃SCH₂, CH₃SCH₂CH₂, CH₃CH₂CH₂CH₂CH₂CH₂CH₂ and CH₃CH₂SCH₂CH₂.

include $(CH_3)_2CHOCH_2C\equiv C$ and $CH_3OCH_2C\equiv C$.

- "Alkylthioalkoxy" denotes alkylthio substitution on alkoxy. Examples of "alkylthioalkoxy" include CH₃SCH₂O and CH₃CH₂SCH₂CH₂O. "Alkylthioalkylthio" denotes alkylthio substitution on alkylthio. Examples of "alkylthioalkylthio" include CH₃SCII₂S and CH₃SCH₂CH₂S. "Alkylsulfinyl" includes both enantiomers of an alkylsulfinyl group. Examples of "alkylsulfinyl" include CH₃S(O), CH₃CH₂S(O),
- CH₃CH₂S(O)₂, CH₃CH₂CH₂S(O)₂, (CH₃)₂CHS(O)₂ and the different butylsulfonyl, pentylsulfonyl, hexylsulfonyl, heptylsulfonyl and octylsulfonyl isomers. "Alkylsulfonylalkyl" denotes alkylsulfonyl substitution on alkyl. Examples of "alkylsulfonylalkyl" include CH₃S(O)₂CH₂, CH₃S(O)₂CH₂CH₂, CH₃CH₂S(O)₂CH₂,

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CH₃CH₂CH₂S(O)₂CH₂ and CH₃CH₂S(O)₂CH₂CH₂. "Alkylthioalkenyl" denotes alkylthio substitution on alkenyl. Examples of "alkylthioalkenyl" include CH₃SCH₂CH=CH and CH₃CH₂SCH=CH. "Alkylthioalkynyl" denotes alkylthio substitution on alkynyl. Examples of "alkylthioalkynyl" include CH₃SCH₂C \equiv C and CH₃CH₂SCH₂CH₂C=C. "Alkenylthio" includes straight-chain and branched alkenylthio moieties. Examples of "alkenylthio" include CH₂=CHCH₂S and CH₂=CHCH₂CH₂S. "Alkenylthioalkyl" denotes alkenylthio substitution on alkyl. Examples of "alkenylthioalkyl" include CH₂=CHCH₂SCH₂ and CH₂=CHCH₂CH₂SCH₂. "Alkynylthio" includes straight-chain and branched alkynylthio moieties. Examples of "alkynylthio" include CH=CCH₂S and CH=CCH₂CH₂S. "Alkynylthioalkyl" denotes alkynylthio substitution on alkyl. Examples of "alkynylthioalkyl" include CH=CCH₂SCH₂ and CH=CCH₂CH₂CH₂SCH₂.

"Cycloalkyl" includes, for example, cyclopropyl, cyclobutyl, cyclopentyl, cyclohexyl, cycloheptyl and cyclooctyl. "(Cycloalkyl)oxyalkyl" denotes cycloalkyl substituted on oxygen which in turn is substituted on alkyl. Examples of "(cycloalkyl)oxyalkyl" include (cyclopentyloxy)methyl and (cyclohexyloxy)methyl. "(Cycloalkyl)thioalkyl" denotes cycloalkyl substituted on sulfur which in turn is substituted on alkyl. Examples of "(cycloalkyl)thioalkyl" include (cyclopentylthio)methyl and (cyclohexylthio)methyl. "Cycloalkenyl" includes groups such as cyclopentenyl and cyclohexenyl as well as groups with more than one double bond such as 1,3- and 1,4-cyclohexadienyl. Examples of "cycloalkylalkyl" include cyclopropylmethyl, cyclopentylethyl, and other cycloalkyl moieties bonded to straight-chain or branched alkyl groups.

"Cyanoalkyl" denotes an alkyl group substituted with one cyano group. Examples of "cyanoalkyl" include $NCCH_2$, $NCCH_2CH_2$ and $CH_3CH(CN)CH_2$. "Nitroalkyl" denotes an alkyl group substituted with one nitro group. Examples of "nitroalkyl" include NO_2CH_2 and $CH_3CH(NO_2)CH_2$.

"Tetrahydropyranyloxyalkynyl" denotes a tetrahydropyranyl group on oxygen which in turn is substituted on an alkynyl group. An example of "tetrahydropyranyloxyalkynyl" is 2-[(tetrahydro-2*H*-pyranyl)oxy]ethynyl.

The term "aromatic ring system" denotes fully unsaturated carbocycles and heterocycles in which the polycyclic ring system is aromatic (where aromatic indicates that the Hückel rule is satisfied for the ring system). The term "aromatic carbocyclic ring system" includes fully aromatic carbocycles and carbocycles in which at least one ring of a polycyclic ring system is aromatic (where aromatic indicates that the Hückel rule is satisfied). The term "nonaromatic carbocyclic ring system" denotes fully saturated carbocycles as well as partially or fully unsaturated carbocycles where the Hückel rule is not satisfied by any of the rings in the ring system. The term "aromatic

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heterocyclic ring system" includes fully aromatic heterocycles and heterocycles in which at least one ring of a polycyclic ring system is aromatic (where aromatic indicates that the Hückel rule is satisfied). Examples of "aromatic heterocyclic ring systems" include furanyl, furazanyl, thienyl, pyrrolyl, pyrazolyl, oxazolyl, oxadiazolyl, imidazolyl, isoxazolyl, thiadiazolyl, thiadiazolyl, isothiazolyl, pyridyl, pyrimidinyl, pyridazinyl, pyrazinyl and triazinyl with said ring attached through any available carbon or nitrogen. For example, when the aromatic heterocyclic ring is furanyl, it can be 2-furanyl or 3-furanyl, for pyrrolyl, the aromatic heterocyclic ring is 1-pyrrolyl, 2-pyrrolyl or 3-pyrrolyl, for pyridyl, the aromatic ring is 2-pyridyl, 3-pyridyl or 4-pyridyl and similarly for other aromatic heterocyclic rings. The term "nonaromatic heterocyclic ring system" denotes fully saturated heterocycles as well as partially or fully unsaturated heterocycles where the Hückel rule is not satisfied by any of the rings in the ring system. The heterocyclic ring systems can be attached through any available

carbon or nitrogen by replacement of a hydrogen on said carbon or nitrogen.

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One skilled in the art will appreciate that not all nitrogen containing heterocycles can form N-oxides since the nitrogen requires an available lone pair for oxidation to the oxide; one skilled in the art will recognize those nitrogen containing heterocycles which can form N-oxides. One skilled in the art will also recognize that tertiary amines can form N-oxides. Synthetic methods for the preparation of N-oxides of heterocycles and tertiary amines are very well known by one skilled in the art including the oxidation of heterocycles and tertiary amines with peroxy acids such as peracetic and m-chloroperbenzoic acid (MCPBA), hydrogen peroxide, alkyl hydroperoxides such as t-butyl hydroperoxide, sodium perborate, and dioxiranes such as dimethydioxirane. These methods for the preparation of N-oxides have been extensively described and reviewed in the literature, see for example: T. L. Gilchrist in Comprehensive Organic Synthesis, vol. 7, pp 748-750, S. V. Ley, Ed., Pergamon Press; M. Tisler and B. Stanovnik in Comprehensive Heterocyclic Chemistry, vol. 3, pp 18-20, A. J. Boulton and A. McKillop, Eds., Pergamon Press; M. R. Grimmett and B. R. T. Keene in Advances in Heterocyclic Chemistry, vol. 43, pp 149-161, A. R. Katritzky, Ed., Academic Press; M. Tisler and B. Stanovnik in *Advances in Heterocyclic Chemistry*, vol. 9, pp 285-291, A. R. Katritzky and A. J. Boulton, Eds., Academic Press; and G. W. H. Cheeseman and E. S. G. Werstiuk in Advances in Heterocyclic Chemistry, vol. 22, pp 390-392, A. R. Katritzky and A. J. Boulton, Eds., Academic Press.

The term "halogen", either alone or in compound words such as "haloalkyl", includes fluorine, chlorine, bromine or iodine. The term "1-2 halogen" indicates that one or two of the available positions for that substituent may be halogen which are independently selected. Further, when used in compound words such as "haloalkyl", said alkyl may be partially or fully substituted with halogen atoms which may be the

same or different. Examples of "haloalkyl" include F₃C, ClCH₂, CF₃CH₂ and CF₃CCl₂. The terms "haloalkenyl", "haloalkenyloxy", "haloalkenylthio", "haloalkenyloxyalkyl", "haloalkynyloxy", "haloalkynyloxyalkyl", "haloalkoxy", "haloalkoxyalkyl", "haloalkylthio", "haloalkylsulfinyl", "haloalkylsulfonyl", and the like, are defined analogously to the term "haloalkyl".

- "haloalkylsulfonyl", and the like, are defined analogously to the term "haloalkyl". Examples of "haloalkenyl" include (Cl)₂C=CHCH₂ and CF₃CH₂CH=CHCH₂. Examples of "haloalkenyloxy" include (Cl)₂C=CHCH₂O and CF₃CH₂CH=CHCH₂O. Examples of "haloalkenylthio" include (Cl)₂C=CHCH₂S and CF₃CH₂CH=CHCH₂S. Examples of "haloalkenyloxyalkyl" include (Cl)₂C=CHCH₂OCH₂ and
- CF₃CH₂CH=CHCH₂OCH₂. Examples of "haloalkynyl" include HC≡CCHCl, CF₃C≡C, CCl₃C≡C and FCH₂C≡CCH₂. Examples of "haloalkynyloxy" include CF₃C≡CCH₂O, CCl₃C≡CCH₂O and FCH₂C≡CCH₂O. Examples of "haloalkynyloxyalkyl" include CCl₃C≡CCH₂OCH₂ and FCH₂C≡CCH₂CH₂OCH₂. Examples of "haloalkoxy" include CF₃O, CCl₃CH₂O, HCF₂CH₂CH₂O and CF₃CH₂O. Examples of "haloalkoxyalkyl"
- include CF₃OCH₂, CCl₃CH₂OCH₂CH₂, HCF₂ CH₂CH₂OCH₂ and CF₃ CH₂OCH₂. Examples of "haloalkylthio" include CCl₃S, CF₃S, CCl₃CH₂S and ClCH₂CH₂CH₂S. Examples of "haloalkylsulfinyl" include CF₃S(O), CCl₃S(O), CF₃CH₂S(O) and CF₃CF₂S(O). Examples of "haloalkylsulfonyl" include CF₃S(O)₂, CCl₃S(O)₂, CCl₃S(O)₂, CF₃CH₂S(O)₂ and CF₃CF₂S(O)₂.
- "Alkylcarbonyl" denotes alkyl substituted carbonyl. Examples of "alkylcarbonyl" include CH₃C(=O) and (CH₃)₂CHC(=O). "Alkoxycarbonyl denotes alkoxy substituted carbonyl. Examples of "alkoxycarbonyl" include CH₃OC(=O) and (CH₃)₂CHOC(=O). "Alkylcarbonylamino" denotes alkylcarbonyl substituted on nitrogen. Examples of "alkylcarbonylamino" include CH₃C(=O)NH and CH₃CH₂C(=O)NH.
- 25 "Alkoxycarbonylamino" denotes alkoxycarbonyl substituted on nitrogen. Examples of "alkoxycarbonylamino" include CH₃OC(=O)NH and CH₃CH₂OC(=O)NH.

Examples of "trialkylsilyl" include $(CH_3)_3Si$ and $(CH_3)_3CSi(CH_3)_2$. "Trialkylsilylalkyl" denotes trialkylsilyl substitution on alkyl. Examples of "trialkylsilylalkyl" include $(CH_3)_3SiCH_2$, and $(CH_3)_3SiCH_2CH_2$.

- "Trialkylsilylalkynyl" denotes trialkylsilyl substitution on alkynyl. Examples of "trialkylsilylalkynyl" include (CH₃)₃SiC≡C and (CH₃CH₂)SiCH₂C≡C.
 "Trialkylsilylalkoxyalkoxy" denotes trialkylsilyl substitution on alkoxy substituted in turn on alkoxy. Examples of "trialkylsilylalkoxyalkoxy" include (CH₃)₃SiCH₂OCH₂O and (CH₃)₃SiCH₂CCH₂OCH₂O.
- The total number of carbon atoms in a substituent group is indicated by the "C_i-C_j" prefix where i and j are numbers from 1 to 11. For example, C₁-C₃ alkylsulfonyl designates methylsulfonyl through propylsulfonyl; C₂ alkoxyalkyl designates CH₃OCH₂; C₃ alkoxyalkyl designates, for example, CH₃CH(OCH₃),

CH₃OCH₂CH₂ or CH₃CH₂OCH₂; and C₄ alkoxyalkyl designates the various isomers of an alkyl group substituted with an alkoxy group containing a total of four carbon atoms, examples including CH₃CH₂CH₂OCH₂ and CH₃CH₂OCH₂CH₂. Examples of "alkylcarbonyl" include C(O)CH₃, C(O)CH₂CH₂CH₃ and C(O)CH(CH₃)₂. Examples of "alkoxycarbonyl" include CH₃OC(=O), CH₃CH₂OC(=O), CH₃CH₂OC(=O), (CH₃)₂CHOC(=O) and the different butoxy- or pentoxycarbonyl isomers. In the above recitations, when a compound of Formula I is comprised of one or more heterocyclic rings, all substituents are attached to these rings through any available carbon or nitrogen by replacement of a hydrogen on said carbon or nitrogen.

When a compound is substituted with a substituent bearing a subscript that indicates the number of said substituents can exceed 1, said substituents (when they exceed 1) are independently selected from the group of defined substituents. Further, when the subscript indicates a range, e.g., $(R)_{i-j}$, then the number of substituents may be selected from the integers between i and j inclusive.

When a group contains a substituent which can be hydrogen, for example R⁴, then, when this substituent is taken as hydrogen, it is recognized that this is equivalent to said group being unsubstituted. When a group is optionally substituted with a substituent, for example with R⁷, then, when the group is not substituted with that substituent, it is recognized that this is equivalent to said group having a hydrogen substituent.

Compounds used in this invention often can exist as one or more stereoisomers. The various stereoisomers include enantiomers, diastereomers, atropisomers and geometric isomers. One skilled in the art will appreciate that one stereoisomer may be more active and/or may exhibit beneficial effects when enriched relative to the other stereoisomer(s) or when separated from the other stereoisomer(s). (See, e.g., U.S. Provisional Patent Application Scrial No. 60/057917 filed September 4, 1997, which is hereby incorporated by reference in its entirety.) Additionally, the skilled artisan knows how to separate, enrich, and/or to selectively prepare said stereoisomers. Accordingly, the present invention comprises compounds selected from Formula I, *N*-oxides and agriculturally suitable salts thereof. The compounds of the invention may be present as a mixture of stereoisomers, individual stereoisomers, or as an optically active form.

The salts of the compounds which may used in the invention include acid-addition salts with inorganic or organic acids such as hydrobromic, hydrochloric, nitric, phosphoric, sulfuric, acetic, butyric, fumaric, lactic, maleic, malonic, oxalic, propionic, salicylic, tartaric, 4-toluenesulfonic or valeric acids. The salts of the compounds which may be used in the invention also include those formed with organic bases (e.g., pyridine, ammonia, or triethylamine) or inorganic bases (e.g., hydrides, hydroxides, or

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carbonates of sodium, potassium, lithium, calcium, magnesium or barium) when the compound contains an acidic group such as a carboxylic acid or phenol.

The fungicidal compositions of this invention, in addition to comprising fungicidally effective amounts of the mixtures of the invention, also optionally comprise at least one of a surfactant, a solid diluent or a liquid diluent. The preferred compositions of the present invention are those which comprise the above preferred component (1) and component (2) compounds.

This invention also relates to a method for controlling plant diseases caused by fungal plant pathogens comprising applying to the plant or portion thereof, or to the plant seed or seedling, a fungicidally effective amount of the compositions of the invention (e.g., as a composition described herein). The preferred methods of use include those involving the above preferred compositions.

The bc | Complex Fungicides

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Compounds of Formula II are known to have a fungicidal mode of action which inhibits the bc_1 complex. Methyl (E)-2-[[6-(2-cyanophenoxy)-4-pyrimidinyl]oxy]- α - (methoxyimino)benzeneacetate is described as a bc_1 complex inhibitor in *Biochemical Society Transactions* 1993, 22, 68S. Methyl (E)- α -(methoxyimino)-2-[(2-methylphenoxy)methyl]benzeneacetate is described as a bc_1 complex inhibitor in *Biochemical Society Transactions* 1993, 22, 64S. (E)-2-[(2,5-

Dimethylphenoxy)methyl]- α -(methoxyimino)-N-methylbenzeneacetamide is described as a bc_1 complex inhibitor in *Biochemistry and Cell Biology* 1995, 85(3), 306-311.

The bc_1 complex is sometimes referred to by other names in the biochemical literature, including complex III of the electron transfer chain, and ubihydroquinone:cytochrome c oxidoreductase. It is uniquely identified by the Enzyme Commission number EC1.10.2.2. The bc_1 complex is described in, for example, *J. Biol. Chem.* 1989, 264, 14543-38; *Methods Enzymol.* 1986, 126, 253-71; and references cited therein.

The Sterol Biosynthesis Inhibitor Fungicides

The class of sterol biosynthesis inhibitors includes DMI and non-DMI compounds, that control fungi by inhibiting enzymes in the sterol biosynthesis pathway. DMI fungicides have a common site of action within the fungal sterol biosynthesis pathway; that is, an inhibition of demethylation at position 14 of lanosterol or 24-methylene dihydrolanosterol, which are precursors to sterols in fungi. Compounds acting at this site are often referred to as demethylase inhibitors, DMI fungicides, or DMIs. The demethylase enzyme is sometimes referred to by other names in the biochemical literature, including cytochrome P-450 (14DM). The demethylase enzyme is described in, for example, *J. Biol. Chem.* 1992, 267, 13175-79 and references cited therein. DMI fungicides fall into several classes: triazoles, imidazoles, pyrimidines,

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piperazines and pyridines. The triazoles includes bromuconazole, cyproconazole, difenoconazole, diniconazole, epoxiconazole, fenbuconazole, fluquinconazole, flusilazole, flutriafol, hexaconazole, ipconazole, metconazole, penconazole, propiconazole, tebuconazole, tetraconazole, triadimefon, triadimenol, triticonazole and uniconazole. The imidazoles include clotrimazole, econazole, imazalil, isoconazole, miconazole and prochloraz. The pyrimidines include fenarimol, nuarimol and triarimol. The piperazines include triforine. The pyridines include buthiobate and pyrifenox. Biochemical investigations have shown that all of the above mentioned fungicides are DMI fungicides as described by K. H. Kuck, et al. in *Modern Selective Fungicides - Properties, Applications and Mechanisms of Action*, Lyr, H., Ed.; Gustav Fischer Verlag: New York, 1995, 205-258.

The DMI fungicides have been grouped together to distinguish them from other sterol biosynthesis inhibitors, such as, the morpholine and piperidine fungicides. The morpholines and piperidines are also sterol biosynthesis inhibitors but have been shown to inhibit later steps in the sterol biosynthesis pathway. The morpholines include aldimorph, dodemorph, fenpropimorph, tridemorph and trimorphamide. The piperidines include fenpropidin. Biochemical investigations have shown that all of the above mentioned morpholine and piperidine fungicides are sterol biosynthesis inhibitor fungicides as described by K. H. Kuck, et al. in *Modern Selective Fungicides* -

Properties, Applications and Mechanisms of Action, Lyr, H., Ed.; Gustav Fischer Verlag: New York, 1995, 185-204.

Synergistic Effects

Fungicides that effectively control plant fungi, particularly wheat powdery mildew (*Erysiphe graminis*) and wheat foot rot (*Pseudocercosporella herpotrichoides*), are in constant demand by growers. Combinations of fungicides are often used to facilitate disease control and to retard resistance development. Mixtures of fungicides may provide significantly better disease control than could be predicted based on the activity of the individual components. This synergism has been described as "the cooperative action of two components of a mixture, such that the total effect is greater or more prolonged than the sum of the effects of the two (or more) taken independently" (see Tames, P. M. L., *Neth. J. Plant Pathology*, (1964), 70, 73-80). It has been demonstrated that compositions containing compounds of Formula Ia and Formula IIa; Formula Ia and flusilazole; Formula Ia and tebuconazole; and Formula Ia, Formula IIa and flusilazole exhibit synergistic effects.

The presence of a synergistic effect between two active ingredients is established with the aid of the Colby equation (see Colby, S. R. In *Calculating Synergistic and Antagonistic Responses of Herbicide Combinations*, Weeds, (1967), 15, 20-22):

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$$p = A + B - \left[\frac{A \times B}{100} \right]$$

Using the methods of the Colby, the presence of a synergistic interaction between two active ingredients is established by first calculating the predicted activity, p, of the mixture based on activities of the two components applied alone. If p is lower than the experimentally established effect, synergism has occurred. In the equation above, A is the fungicidal activity in percentage control of one component applied alone at rate x. The B term is the fungicidal activity in percentage control of the second component applied at rate y. The equation estimates p, the fungicidal activity of the mixture of A at rate x with B at rate y if their effects are strictly additive and no interaction has occurred.

The Colby equation for a three way mixture is

$$p = A + B + C + \frac{A \times B \times C}{10000} - \frac{(A \times B) + (A \times C) + (B \times C)}{100}$$

In this application, fungicidal activities provided by compositions of Formula Ia, Formula IIa, flusilazole and tebuconazole alone are compared with that of compositions of the compounds of Formula Ia and Formula IIa together, compositions of the compounds of Formula Ia and flusilazole together, compositions of the compounds of Formula Ia and tebuconazole together, and compositions of the compounds of Formula Ia, Formula IIa and flusilazole together. Based on the description of synergism developed by Colby, compositions of the present invention are considered to be synergistically useful. Accordingly, this invention provides an improved method of combating fungi, such as wheat powdery mildew (*Erysiphe graminis*), wheat leaf rust (*Puccinia recondita*), wheat foot rot (*Pseudocercosporella herpotrichoides*) and/or wheat glume blotch (*Septoria nodorum*) in crops, especially cereals.

Compositions are provided in accordance with this invention which comprise proportions of component (1) and component (2) which are especially useful for controlling particular fungal diseases. For example, the compositions of this invention include those wherein the mole ratio of component (1) to component (2) is from about 30:1 to 1:30. These compositions are considered especially useful for controlling wheat powdery mildew (*Erysiphe graminis*), wheat leaf rust (*Puccinia recondita*), wheat foot rot (*Pseudocercosporella herpotrichoides*) and/or wheat glume blotch (*Septoria nodorum*). Preferred component (1) compounds for these compositions include 6-bromo-3-propyl-2-propyloxy-4(3*H*)-quinazolinone; 6,8-diiodo-3-propyl-2-propyloxy-4(3*H*)-quinazolinone; with

6-iodo-3-propyl-2-propyloxy-4(3H)-quinazolinone being particularly preferred.

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Preferred component (2) compounds for these compositions include 2,4-dihydro-5-methoxy-2-methyl-4-[2-[[[[1-[3-

(trifluoromethyl)phenyl]ethylidene]amino]oxy]methyl]phenyl]-3*H*-1,2,4-triazol-3-one, kresoxim-methyl, azoxystrobin, (*E*)-2-(methoxyimino)-*N*-methyl-2-(2-

phenoxyphenyl)acetamide, flusilazole, epoxiconazole, fenpropimorph, propiconazole and tebuconazole; with 2,4-dihydro-5-methoxy-2-methyl-4-[2-[[[[1-[3-(trifluoromethyl)phenyl]ethylidene]amino]oxy]methyl]phenyl]-3*H*-1,2,4-triazol-3-one, fenpropimorph, flusilazole, epoxiconazole and propiconazole being particularly preferred. Preferably, the mole ratio of component (1) to component (2) for these compositions is from about 4:1 to 1:4. Example compositions of this type include compositions comprising 6-iodo-3-propyl-2-propyloxy-4(3*H*)-quinazolinone and 2,4-

dihydro-5-methoxy-2-methyl-4-[2-[[[[1-[3-

- (trifluoromethyl)phenyl]ethylidene]amino]oxy]methyl]phenyl]-3*H*-1,2,4-triazol-3-one in a mole ratio of the compound of Formula Ia to the compound of Formula IIa of from about 4:1 to 1:4, compositions comprising
- 6-iodo-3-propyl-2-propyloxy-4(3*H*)-quinazolinone and fenpropimorph in a mole ratio of the compound of Formula Ia to fenpropimorph of from about 1:1 to 1:10, compositions comprising 6-iodo-3-propyl-2-propyloxy-4(3*H*)-quinazolinone and flusilazole in a mole ratio of the compound of Formula Ia to flusilazole of from about 15:1 to 1:15,
- compositions comprising 6-iodo-3-propyl-2-propyloxy-4(3*H*)-quinazolinone and tebuconazole in a mole ratio of the compound of Formula Ia to tebuconazole of from about 30:1 to 1:30, and compositions comprising
 - 6-iodo-3-propyl-2-propyloxy-4(3*H*)-quinazolinone; 2,4-dihydro-5-methoxy-2-methyl-4-[2-[[[[1-[3-(trifluoromethyl)phenyl]ethylidene]amino]oxy]methyl]phenyl]-3*H*-1,2,4-
- triazol-3-one; and flusilazole in a mole ratio of the compound of Formula Ia to the compound of Formula IIa plus flusilazole of from about 4:1 to 1:4 and the compound of Formula IIa to flusilazole has a mole ratio of from about 4:1 to 1:4.

This invention also provides a process for controlling at least one plant disease selected from wheat powdery mildew, wheat leaf rust, wheat foot rot, and wheat glume blotch which comprises applying to the plant or portion thereof to be protected, or to the plant seed or seedling to be protected, an effective amount of a fungicidal combination including component (1) and component (2) wherein the mole ratio of component (1) to component (2) is from about 30:1 to 1:30 (preferably from about 4:1 to 1:4).

Component (1) can, for example, be applied at a rate of 0.2 g/ha or more. Typically component (1) is applied at a rate of 100 g/ha. Component (2) may be applied simultaneously (e.g., in the form of a composition comprising component (1) and component (2) can

be applied separately in an appropriate mole ratio (e.g., as a tank mix).

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Compositions wherein component (2) is selected from the group consisting of 2,4-dihydro-5-methoxy-2-methyl-4-[2-[[[[1-[3-(trifluoromethyl)phenyl]ethylidene]amino]oxy]methyl]phenyl]-3H-1,2,4-triazol-3-one, kresoxim-methyl, azoxystrobin, (E)-2-(methoxyimino)-N-methyl-2-(2-

- phenoxyphenyl)acetamide, flusilazole, epoxiconazole, fenpropimorph, propiconazole and tebuconazole; and wherein the mole ratio of component (1) to component (2) is from about 4:1 to 1:10 are considered especially useful for controlling wheat powdery mildew. Preferred component (1) compounds for these compositions include 6-bromo-3-propyl-2-propyloxy-4(3*H*)-quinazolinone;
- 6,8-diiodo-3-propyl-2-propyloxy-4(3*H*)-quinazolinone; and 6-iodo-3-propyl-2-propyloxy-4(3*H*)-quinazolinone; with 6-iodo-3-propyl-2-propyloxy-4(3*H*)-quinazolinone being particularly preferred. Preferred component (2) compounds for these compositions include 2,4-dihydro-5-methoxy-2-methyl-4-[2-[[[1-[3-
- (trifluoromethyl)phenyl]ethylidene]amino]oxy]methyl]phenyl]-3*H*-1,2,4-triazol-3-one, fenpropimorph, flusilazole, epoxiconazole and propiconazole. Example compositions of this type include compositions comprising 6-iodo-3-propyl-2-propyloxy-4(3*H*)-quinazolinone and 2,4-dihydro-5-methoxy-2-methyl-4-[2-[[[[1-[3-(trifluoromethyl)phenyl]ethylidene]amino]oxy]methyl]phenyl]-3*H*-
- 1,2,4-triazol-3-one in a mole ratio of the compound of Formula Ia to the compound of Formula IIa of from about 4:1 to 1:4, compositions comprising
 6-iodo-3-propyl-2-propyloxy-4(3H)-quinazolinone and fenpropimorph in a mole ratio of the compound of Formula Ia to fenpropimorph of from about 1:1 to 1:10, compositions comprising 6-iodo-3-propyl-2-propyloxy-4(3H)-quinazolinone and flusilazole in a mole
- ratio of the compound of Formula Ia to flusilazole of from about 4:1 to 1:4, and compositions comprising 6-iodo-3-propyl-2-propyloxy-4(3*H*)-quinazolinone; 2,4-dihydro-5-methoxy-2-methyl-4-[2-[[[[1-[3-
 - (trifluoromethyl)phenyl]ethylidene]amino]oxy]methyl]phenyl]-3*H*-1,2,4-triazol-3-one; and flusilazole in a mole ratio of the compound of Formula Ia to the total of the compound of Formula IIa plus flusilazole of from about 4:1 to 1:4 and a mole ratio of

the compound of Formula IIa to flusilazole of from about 4:1 to 1:4.

This invention also provides a process for controlling wheat powdery mildew which comprises applying to the plant or portion thereof to be protected, or to the plant seed or seedling to be protected, an effective amount of a fungicidal combination including component (1) and component (2) wherein the mole ratio of component (1) to component (2) is from about 4:1 to 1:10 (preferably from about 4:1 to 1:4). Component (1) can, for example, be applied at a rate of 0.2 g/ha or more. Typically component (1) is applied at a rate of 100 g/ha. Component (2) may be applied

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simultaneously (e.g., in the form of a composition comprising component (1) and component (2) in an appropriate mole ratio); or component (1) and component (2) can be applied separately in an appropriate mole ratio (e.g., as a tank mix).

Compositions wherein component (2) is selected from the group consisting of 2,4-5 dihydro-5-methoxy-2-methyl-4-[2-[[[]1-[3-(trifluoromethyl)phenyl lethylidene lamino loxy methyl lphenyl 3H-1.2.4-triazol-3-one, kresoxim-methyl, azoxystrobin, (E)-2-(methoxyimino)-N-methyl-2-(2phenoxyphenyl)acetamide, flusilazole, epoxiconazole, fenpropimorph, propiconazole and tebuconazole; and wherein the mole ratio of component (1) to component (2) is 10 from about 30:1 to 1:30 are considered especially useful for controlling wheat leaf rust. Preferred component (1) compounds for these compositions include 6-bromo-3-propyl-2-propyloxy-4(3H)-quinazolinone; 6,8-diiodo-3-propyl-2-propyloxy-4(3H)-quinazolinone; and 6-iodo-3-propyl-2-propyloxy-4(3H)-quinazolinone; with 15 6-iodo-3-propyl-2-propyloxy-4(3H)-quinazolinone being particularly preferred being particularly preferred. Preferred component (2) compounds for these compositions include 2,4-dihydro-5-methoxy-2-methyl-4-[2-[[[[1-[3-(trifluoromethyl)phenyl]ethylidene|amino|oxy|methyl]phenyl]-3H-1,2,4-triazol-3-one, epoxiconazole, flusilazole and tebuconazole; with flusilazole and epoxiconazole being 20 particularly preferred. Preferably, the mole ratio of component (1) to component (2) for these compositions if from about 4:1 to 1:4. Example compositions of this type include compositions comprising 6-iodo-3-propyl-2-propyloxy-4(3H)-quinazolinone and flusilazole in a mole ratio of the compound of Formula Ia to flusilazole of from about

6-iodo-3-propyl-2-propyloxy-4(3*H*)-quinazolinone and tebuconazole in a mole ratio of the compound of Formula Ia to tebuconazole from about 30:1 to 1:30.

15:1 to 1:15, compositions comprising

This invention also provides a process for controlling wheat leaf rust which comprises applying to the plant or portion thereof to be protected, or to the plant seed or seedling to be protected, an effective amount of a fungicidal combination including component (1) and component (2) wherein the mole ratio of component (1) to component (2) is from about 30:1 to 1:30 (preferably from about 4:1 to 1:4). Component (2) can, for example, be applied at a rate of 12.5 g/ha or more. Typically component (2) is applied at a rate of 160 g/ha. Component (1) may be applied simultaneously (e.g., in the form of a composition comprising component (1) and component (2) can be applied separately in an appropriate mole ratio (e.g., as a tank mix).

Compositions wherein component (2) is selected from the group consisting of 2,4-dihydro-5-methoxy-2-methyl-4-[2-[[[[1-[3-

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(trifluoromethyl)phenyl]ethylidene]amino]oxy]methyl]phenyl]-3*H*-1,2,4-triazol-3-one, kresoxim-methyl, azoxystrobin, (*E*)-2-(methoxyimino)-*N*-methyl-2-(2-phenoxyphenyl)acetamide, flusilazole, epoxiconazole, fenpropimorph, propiconazole and tebuconazole; and wherein the mole ratio of component (1) to component (2) is from about 15:1 to 1:15 are considered especially useful for controlling wheat foot rot. Preferred component (1) compounds for these compositions include 6-bromo-3-propyl-2-propyloxy-4(3*H*)-quinazolinone; 6,8-diiodo-3-propyl-2-propyloxy-4(3*H*)-quinazolinone; with 6-iodo-3-propyl-2-propyloxy-4(3*H*)-quinazolinone being particularly preferred being particularly preferred. A particularly preferred component (2) compound for these

6-iodo-3-propyl-2-propyloxy-4(3*H*)-quinazolinone being particularly preferred being particularly preferred. A particularly preferred component (2) compound for these compositions is flusilazole. Preferably, the mole ratio of component (1) to component (2) for these compositions if from about 4:1 to 1:4. Example compositions of this type include compositions comprising 6-iodo-3-propyl-2-propyloxy-4(3*H*)-quinazolinone and flusilazole in a mole ratio of the compound of Formula la to flusilazole of from about 15:1 to 1:15.

This invention also provides a process for controlling wheat foot rot which comprises applying to the plant or portion thereof to be protected, or to the plant seed or seedling to be protected, an effective amount of a fungicidal combination including component (1) and component (2) wherein the mole ratio of component (1) to component (2) is from about 15:1 to 1:15 (preferably from about 4:1 to 1:4). Component (1) can, for example, be applied at a rate of 5 g/ha or more. Typically component (1) is applied at a rate of 100 g/ha. Component (2) may be applied simultaneously (e.g., in the form of a composition comprising component (1) and component (2) in an appropriate mole ratio); or component (1) and component (2) can be applied separately in an appropriate mole ratio (e.g., as a tank mix).

Compositions wherein component (2) is selected from the group consisting of 2,4-dihydro-5-methoxy-2-methyl-4-[2-[[[1-[3-(trifluoromethyl)phenyl]ethylidene]amino]oxy]methyl]phenyl]-3*H*-1,2,4-triazol-3-one, kresoxim-methyl, azoxystrobin, (*E*)-2-(methoxyimino)-*N*-methyl-2-(2-phenoxyphenyl)acetamide, flusilazole, epoxiconazole, fenpropimorph, propiconazole and tebuconazole; and wherein the mole ratio of component (1) to component (2) is from about 30:1 to 1:30 are considered especially useful for controlling wheat glume blotch. Preferred component (1) compounds for these compositions include 6-bromo-3-propyl-2-propyloxy-4(3*H*)-quinazolinone;

6-iodo-3-propyl-2-propyloxy-4(3*H*)-quinazolinone; with 6-iodo-3-propyl-2-propyloxy-4(3*H*)-quinazolinone being particularly preferred being

6,8-diiodo-3-propyl-2-propyloxy-4(3H)-quinazolinone; and

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particularly preferred. Preferred component (2) compounds for these compositions include flusilazole and tebuconazole; with flusilazole being particularly preferred. Preferably, the mole ratio of component (1) to component (2) for these compositions if from about 4:1 to 1:4. Example compositions of this type include compositions comprising 6-iodo-3-propyl-2-propyloxy-4(3*H*)-quinazolinone and tebuconazole in a mole ratio of the compound of Formula Ia to tebuconazole from about 30:1 to 1:30

This invention also provide a process for controlling wheat glume blotch which comprises applying to the plant or portion thereof to be protected, or to the plant seed or seedling to be protected, an effective amount of a fungicidal combination including component (1) and component (2) wherein the mole ratio of component (1) to component (2) is from about 30:1 to 1:30 (preferably from about 4:1 to 1:4). Component (2) can, for example, be applied at 12.5 g/ha or more. Typically component (2) is applied at a rate of 160 g/ha. Component (1) may be applied simultaneously (e.g., in the form of a composition comprising component (1) and component (2) in an appropriate mole ratio); or component (1) and component (2) can be applied separately in an appropriate mole ratio, e.g., as a tank mix.

Synthesis of Compounds of Formula I

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The compounds of Formula I where D is O can be prepared as described in International Patent Application WO 94/26722 and as shown in Scheme 1.

An anthranilic acid (2-aminobenzoic acid) of Formula 1 is condensed with an isothiocyanate of Formula R¹-NCS to form the 2-thioquinazolinedione of Formula 2. This condensation is preferably performed in the presence of a base such as triethylamine. S-Methylation of this compound affords the 2-methylthio-4(3H)-quinazolinone of Formula 3.

For the introduction of the R²O group, the 2-methylthio-4(3*H*)-quinazolinone of Formula 3 is treated with a mixture of a base, for example sodium hydride, in R²OH solvent. The reaction mixture is stirred at a temperature from about 0 °C to 120 °C for 1-120 hours. The desired 2-R²O-4(3H)-quinazolinone can be isolated from the reaction mixture by extraction into a water-immiscible solvent, and purified by chromatography or recrystallization. Similar synthetic procedures are described in U.S. 3,755,582, incorporated herein by reference.

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Scheme 1

Anthranilic acids of Formula 1 are known or can be prepared by known methods. For example see, March, J. *Advanced Organic Chemistry*; 3rd ed., John Wiley:

New York, (1985), p 983. The isothiocyanates of Formula R¹-NCS can be prepared from the corresponding amine by treatment with thiophosgene as known in the art. For example, see *J. Heterocycl. Chem.*, (1990), *27*, 407.

6-lodo-3-propyl-2-propyloxy-4(3H)-quinazolinone, Formula Ia, is a compound of Formula I where D is O, R^1 is propyl, R^2 is propyl, R^3 is iodine fixed in the 6 position and R^4 is hydrogen.

Synthesis of Compounds of Formula II

Compounds of Formula II where Q is Q-1 and X is OR¹¹ can be prepared as described in International Patent Application WO 95/14009 by treating a compound of Formula 4 with an appropriate alkyl transfer reagent in an inert solvent with or without additional acidic or basic reagents or other reagents (Scheme 2). Suitable solvents are selected from the group consisting of polar aprotic solvents such as acetonitrile, dimethylformamide or dimethylsulfoxide; ethers such as tetrahydrofuran, dimethoxyethane, or diethyl ether; ketones such as acetone or 2-butanone; hydrocarbons such as toluene or benzene; and halocarbons such as dichloromethane or chloroform.

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Scheme 2

HO
$$\frac{E}{A}$$
 $\frac{Y}{N}$ $\frac{E}{A}$ $\frac{Y}{N}$ $\frac{Z}{A}$ $\frac{E}{N}$ $\frac{Y}{N}$ $\frac{Z}{N}$ \frac

Method 1: $V-CH=N_2$ ($V = H \text{ or } (CH_3)_3Si$)

Method 2: Cl₃C OR¹¹; Lewis acid

Method 3: $(R^{11})_3O^+BF_4^-$

Method 4: $(R^{11})_2SO_4$; $R^{11}OSO_2V$; or R^{11} -hal; optional base (hal = F, Cl, Br, or I) $(V = C_1-C_6 \text{ alkyl}, C_1-C_6 \text{ haloalkyl})$

For example, compounds of Formula II where Q is Q-1 and X is OR^{11} can be prepared by the action of diazoalkane reagents of Formula 5 such as diazomethane (V = H) or trimethylsilyldiazomethane (V = $(CH_3)_3Si$) on of Formula 4 (Method 1). Use of trimethylsilyldiazomethane requires a protic cosolvent such as methanol. For examples of these procedures, see *Chem. Pharm. Bull.*, (1984), 32, 3759.

As indicated in Method 2, compounds of Formula II where Q is Q-1 and X is Q-1 can also be prepared by contacting compounds of Formula 4 with alkyl

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trichloroacetimidates of Formula 6 and a Lewis acid catalyst. Suitable Lewis acids include trimethylsilyl triflate and tetrafluoroboric acid. The alkyl trichloroacetimidates can be prepared from the appropriate alcohol and trichloroacetonitrile as described in the literature (J. Danklmaier and H. Hönig, *Synth. Commun.*, (1990), 20, 203).

Compounds of Formula II where Q is Q-1 and X is OR¹¹ can also be prepared from compounds of Formula 4 by treatment with a trialkyloxonium tetrafluoroborate (i.e., Meerwein's salt) of Formula 7 (Method 3). The use of trialkyloxonium salts as powerful alkylating agents is well known in the art (see U. Schöllkopf, U. Groth, C. Deng, *Angew. Chem., Int. Ed. Engl.*, (1981), 20, 798).

Other alkylating agents which can convert compounds of Formula 4 to compounds of Formula II where Q is Q-1 and X is OR¹¹ are dialkyl sulfates such as dimethyl sulfate, haloalkyl sulfonates such as methyl trifluoromethanesulfonate, and alkyl halides such as iodomethane and propargyl bromide (Method 4). These alkylations can be conducted with or without additional base. Appropriate bases include alkali metal alkoxides such as potassium *tert*-butoxide, inorganic bases such as sodium hydride and potassium carbonate, or tertiary amines such as triethylamine, pyridine, 1,8-diazabicyclo[5.4.0]undec-7-ene (DBU), and triethylenediamine. See R. E. Benson, T. L. Cairns, *J. Am. Chem. Soc.*, (1948), 70, 2115 for alkylation examples using agents of this type.

Compounds of Formula II where Q is Q-1 and X is OR¹¹ can also be prepared by reaction of Formula 8 compounds with alkali metal alkoxides (R¹¹O-M⁺) (Scheme 3). The leaving group Lg¹ in the amides of Formula 8 are any group known in the art to undergo a displacement reaction of this type. Examples of suitable leaving groups include chlorine, bromine, and sulfonyl and sulfonate groups. Examples of suitable inert solvents are dimethylformamide or dimethylsulfoxide.

Scheme 3

Lg
$$\frac{E}{G}$$
 W

R110 $\frac{G}{G}$ W

R12

8 Lg $\frac{1}{G}$ = Cl, Br, $-SO_2V$ or $-OSO_2V$ II

 $V = C_1 - C_6$ alkyl or $C_1 - C_6$ haloalkyl

M = K or Na

2,4-Dihydro-5-methoxy-2-methyl-4-[2-[[[[1-[3-(trifluoromethyl)phenyl]ethylidene]amino]oxy]methyl]phenyl]-3*H*-1,2,4-triazol-3-one, Formula IIa, is a compound of Formula II where Q is Q-1, E is 1,2-phenylene, A is N, G is N with the floating double bond attached to A, W is O, X is OR¹¹, R¹¹ is CH₃, R¹² is CH₃, Y is -CHR²⁵O-N=C(R¹⁷)-, R¹⁷ is CH₃, R²⁵ is H, Z is phenyl substituted with R¹⁹ and R¹⁹ is CF₃ fixed in the 3 position.

Azoxystrobin, Formula IIb, is a compound of Formula II where Q is Q-2, E is 1,2 phenylene, W^1 is O, X^1 is C, Y is -O-, Z is 4,6-pyrimidinyl substituted with R^{19} , R^{19} is phenoxy substituted with R^{21} and R^{21} is cyano fixed in the 2 position.

Kresoxim-methyl, Formula IIc, is a compound of Formula II where Q is Q-2, E is 1,2-phenylene, W¹ is O, X¹ is N, Y is -CHR²⁵O-, R²⁵ is H, Z is phenyl substituted with R¹⁹ and R¹⁹ is CH₃ fixed in the 2 position.

(*E*)-2-(methoxyimino)-*N*-methyl-2-(2-phenoxyphenyl)acetamide, Formula IId, is a compound of Formula II where Q is Q-2, E is 1,2-phenylene, W¹ is NH, X¹ is N, Y is -O-, Z is phenyl substituted with R¹⁹ wherein R¹⁹ is H.

Without further elaboration, it is believed that one skilled in the art using the preceding description can utilize the present invention to its fullest extent. The following Examples are, therefore, to be construed as merely illustrative.

20 Formulation/Utility

The fungicidal compositions of the present invention comprise an effective amount of a mixture of the compounds(s) of component (1) (e.g., 6-iodo-3-propyl-2-propyloxy-4(3*H*)-quinazolinone) and the compound(s) of component (2) (e.g., 2,4-dihydro-5-methoxy-2-methyl-4-[2-[[[[1-[3-

(trifluoromethyl)phenyl]ethylidene]amino]oxy]methyl]phenyl]-3H-1,2,4-triazol-3-one, kresoxim-methyl, azoxystrobin, (E)-2-(methoxyimino)-N-methyl-2-(2-phenoxyphenyl)acetamide, flusilazole, epoxiconazole, fenpropimorph, propiconazole and/or tebuconazole). The mixtures of this invention will typically be used as a formulation or composition with an agriculturally suitable carrier comprising at least one of a liquid diluent, a solid diluent or a surfactant. The formulation or composition

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ingredients are selected to be consistent with the physical properties of the active ingredient, mode of application and environmental factors such as soil type, moisture and temperature. Useful formulations include liquids such as solutions (including emulsifiable concentrates), suspensions, emulsions (including microemulsions and/or suspoemulsions) and the like which optionally can be thickened into gels. Useful formulations further include solids such as dusts, powders, granules, pellets, tablets, films, and the like which can be water-dispersible ("wettable") or water-soluble. Active ingredient can be (micro)encapsulated and further formed into a suspension or solid formulation; alternatively the entire formulation of active ingredient can be encapsulated (or "overcoated"). Encapsulation can control or delay release of the active ingredient. Sprayable formulations can be extended in suitable media and used at spray volumes from about one to several hundred liters per hectare. High-strength compositions are primarily used as intermediates for further formulation.

The formulations will typically contain effective amounts of active ingredients, diluent and surfactant within the following approximate ranges which add up to 100 percent by weight.

		Weight Percent	
	Active Ingredients	Diluent	Surfactant
Water-Dispersible and Water-soluble Granules, Tablets and Powders.	5–90	0–94	1–15
Suspensions, Emulsions, Solutions (including Emulsifiable Concentrates)	550	40–95	0-15
Dusts Granules and Pellets	1–25 0.01–99	70–99 5–99.99	0-5 0-15
High Strength Compositions	90-99	0-10	0-2

Typical solid diluents are described in Watkins, et al., Handbook of Insecticide Dust Diluents and Carriers, 2nd Ed., Dorland Books, Caldwell, New Jersey. Typical liquid diluents are described in Marsden, Solvents Guide, 2nd Ed., Interscience, New York, 1950. McCutcheon's Detergents and Emulsifiers Annual, Allured Publ. Corp., Ridgewood, New Jersey, as well as Sisely and Wood, Encyclopedia of Surface Active Agents, Chemical Publ. Co., Inc., New York, 1964, list surfactants and recommended uses. All formulations can contain minor amounts of additives to reduce foam, caking, corrosion, microbiological growth and the like, or thickeners to increase viscosity.

Surfactants include, for example, polyethoxylated alcohols, polyethoxylated alkylphenols, polyethoxylated sorbitan fatty acid esters, dialkyl sulfosuccinates, alkyl sulfates, alkylbenzene sulfonates, organosilicones, *N*,*N*-dialkyltaurates, lignin

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sulfonates, naphthalene sulfonate formaldehyde condensates, polycarboxylates, and polyoxyethylene/polyoxypropylene block copolymers. Solid diluents include, for example, clays such as bentonite, montmorillonite, attapulgite and kaolin, starch, sugar, silica, tale, diatomaceous earth, urea, calcium carbonate, sodium carbonate and bicarbonate, and sodium sulfate. Liquid diluents include, for example, water, *N*,*N*-dimethylformamide, dimethyl sulfoxide, *N*-alkylpyrrolidone, ethylene glycol, polypropylene glycol, paraffins, alkylbenzenes, alkylnaphthalenes, oils of olive, castor, linseed, tung, sesame, corn, peanut, cotton-seed, soybean, rape-seed and coconut, fatty acid esters, ketones such as cyclohexanone, 2-heptanone, isophorone and 4-hydroxy-4-methyl-2-pentanone, and alcohols such as methanol, cyclohexanol, decanol and tetrahydrofurfuryl alcohol.

Solutions, including emulsifiable concentrates, can be prepared by simply mixing the ingredients. Dusts and powders can be prepared by blending and, usually, grinding as in a hammer mill or fluid-energy mill. Suspensions are usually prepared by wet-milling; see, for example, U.S. 3,060,084. Granules and pellets can be prepared by spraying the active material upon preformed granular carriers or by agglomeration techniques. See Browning, "Agglomeration", *Chemical Engineering*, December 4, 1967, pp 147-48, *Perry's Chemical Engineer's Handbook*, 4th Ed., McGraw-Hill, New York, 1963, pages 8-57 and following, and International Patent Publication WO 91/13546. Pellets can be prepared as described in U.S. Patent No. 4 172 714

91/13546. Pellets can be prepared as described in U.S. Patent No. 4,172,714.
Water-dispersible and water-soluble granules can be prepared as taught in
U.S. Patent No. 4,144,050, U.S. Patent No. 3,920,442 and German Patent Application
DE 3,246,493. Tablets can be prepared as taught in U.S. Patent No. 5,180,587,
U.S. Patent No. 5,232,701 and U.S. Patent No. 5,208,030. Films can be prepared as
taught in Great Britain Patent Application GB 2,095,558 and U.S. Patent No. 3,299,566.

For further information regarding the art of formulation, see U.S. Patent No. 3,235,361, Col. 6, line 16 through Col. 7, line 19 and Examples 10-41; U.S. Patent No. 3,309,192, Col. 5, line 43 through Col. 7, line 62 and Examples 8, 12, 15, 39, 41, 52, 53, 58, 132, 138-140, 162-164, 166, 167 and 169-182;

30 U.S. Patent No. 2,891,855, Col. 3, line 66 through Col. 5, line 17 and Examples 1-4; Klingman, Weed Control as a Science, John Wiley and Sons, Inc., New York, 1961, pp 81-96; and Hance et al., Weed Control Handbook, 8th Ed., Blackwell Scientific Publications, Oxford, 1989.

In the following Examples, all percentages are by weight and all formulations are prepared in conventional ways.

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Example A

	Wettable Powder	
	Active ingredient(s)	65.0%
	dodecylphenol polyethylene glycol ether	2.0%
5	sodium ligninsulfonate	4.0%
	sodium silicoaluminate	6.0%
	montmorillonite (calcined)	23.0%.
	Example B	
	Granule	
10	Active ingredient(s)	10.0%
	attapulgite granules (low volatile matter,	
	0.71/0.30 mm; U.S.S. No. 25-50 sieves)	90.0%.
	Example C	
	Extruded Pellet	
15	Active ingredient(s)	25.0%
	anhydrous sodium sulfate	10.0%
	crude calcium ligninsulfonate	5.0%
	sodium alkylnaphthalenesulfonate	1.0%
	calcium/magnesium bentonite	59.0%.
20	Example D	
	Emulsifiable Concentrate	
	Active ingredient(s)	20.0%
	blend of oil soluble sulfonates	
	and polyoxyethylene ethers	10.0%
25	isophorone	70.0%.

The compounds of this invention are useful as plant disease control agents. The present invention therefore further comprises a method for controlling plant diseases caused by fungal plant pathogens comprising applying to the plant or portion thereof to be protected, or to the plant seed or seedling to be protected, an effective amount of a compound of the invention or a fungicidal composition containing said compound. The compounds and compositions of this invention provide control of diseases caused by a broad spectrum of fungal plant pathogens in the Basidiomycete, Ascomycete, Oomycete and Deuteromycete classes. They are effective in controlling a broad spectrum of plant diseases, particularly foliar pathogens of ornamental, vegetable, field, cereal, and fruit crops. These pathogens include *Plasmopara viticola*, *Phytophthora infestans*, *Peronospora tabacina*, *Pseudoperonospora cubensis*, *Pythium aphanidermatum*, *Alternaria brassicae*, *Septoria nodorum*, *Septoria tritici*, *Cercosporidium personatum*, *Cercospora arachidicola*, *Pseudocercosporella herpotrichoides*, *Cercospora beticola*,

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Botrytis cinerea, Monilinia fructicola, Pyricularia oryzae, Podosphaera leucotricha, Venturia inaequalis, Erysiphe graminis, Uncinula necatur, Puccinia recondita, Puccinia graminis, Hemileia vastatrix, Puccinia striiformis, Puccinia arachidis, Rhizoctonia solani, Sphaerotheca fuliginea, Fusarium oxysporum, Verticillium dahliae, Pythium aphanidermatum, Phytophthora megasperma, Sclerotinia sclerotiorum, Sclerotium rolfsii, Erysiphe polygoni, Pyrenophora teres, Gaeumannomyces graminis, Rynchosporium secalis, Fusarium roseum, Bremia lactucae and other generea and species closely related to these pathogens.

Mixtures of this invention can be further mixed with one or more other 10 insecticides, fungicides, nematocides, bactericides, acaricides, growth regulators, chemosterilants, semiochemicals, repellents, attractants, pheromones, feeding stimulants or other biologically active compounds to form a multi-component pesticide giving an even broader spectrum of agricultural protection. Examples of such agricultural protectants with which compounds of this invention can be formulated are: insecticides 15 such as abamectin, acephate, azinphos-methyl, bifenthrin, buprofezin, carbofuran, chlorfenapyr, chlorpyrifos, chlorpyrifos-methyl, cyfluthrin, beta-cyfluthrin, cyhalothrin, lambda-cyhalothrin, deltamethrin, diafenthiuron, diazinon, diflubenzuron, dimethoate, esfenvalerate, fenoxycarb, fenpropathrin, fenvalerate, fipronil, flucythrinate, tau-fluvalinate, fonophos, imidacloprid, isofenphos, malathion, metaldehyde, 20 methamidophos, methidathion, methomyl, methoprene, methoxychlor, methyl 7-chloro-2,5-dihydro-2-[[N-(methoxycarbonyl)-N-[4-(trifluoromethoxy)phenyl]amino]carbonyl]indeno[1,2-e][1,3,4]oxadiazine-4a(3H)carboxylate (DPX-JW062), monocrotophos, oxamyl, parathion, parathion-methyl, permethrin, phorate, phosalone, phosmet, phosphamidon, pirimicarb, profenofos, 25 rotenone, sulprofos, tebufenozide, tefluthrin, terbufos, tetrachlorvinphos, thiodicarb, tralomethrin, trichlorfon and triflumuron; fungicides such as benomyl, blasticidin-S, Bordeaux mixture (tribasic copper sulfate), captafol, captan, carbendazim, chloroneb, chlorothalonil, copper oxychloride, copper salts, cymoxanil, cyprodinil (CGA 219417), diclomezine, dicloran, dimethomorph, dodine, edifenphos, famoxadone, fenpiclonil, 30 fluazinam, flutolanil, folpet, fosetyl-aluminum, furalaxyl, iprobenfos, iprodione, isoprothiolane, kasugamycin, mancozeb, maneb, mepronil, metalaxyl, S-methyl 7-benzothiazolecarbothioate (CGA 245704), myclobutanil, neo-asozin (ferric methanearsonate), oxadixyl, pencycuron, probenazole, prochloraz, pyrifenox, pyroquilon, quinoxyfen, spiroxamine (KWG4168), sulfur, thiabendazole, 35 thiophanate-methyl, thiram, tricyclazole, validamycin and vinclozolin; nematocides such as aldoxycarb and fenamiphos; bactericides such as streptomycin; acaricides such

as amitraz, chinomethionat, chlorobenzilate, cyhexatin, dicofol, dienochlor, etoxazole, fenazaquin, fenbutatin oxide, fenpropathrin, fenpyroximate, hexythiazox, propargite,

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pyridaben and tebufenpyrad; and biological agents such as *Bacillus thuringiensis*, *Bacillus thuringiensis* delta endotoxin, baculovirus, and entomopathogenic bacteria, virus and fungi.

In certain instances, combinations with other fungicides having a similar spectrum of control but a different mode of action will be particularly advantageous for resistance management.

Plant disease control is ordinarily accomplished by applying an effective amount of the composition of this invention either pre— or post—infection, to the portion of the plant to be protected such as the roots, stems, foliage, fruit, seeds, tubers or bulbs, or to the media (soil or sand) in which the plants to be protected are growing. The composition can also be applied to the seed to protect the seed and seedling.

Rates of application for this composition can be influenced by many factors of the environment and should be determined under actual use conditions. Foliage can normally be protected when treated at a rate of from less than 1 g/ha to 5,000 g/ha of aggregate active ingredient. Aggregate active ingredient is defined as the total combined weight of active ingredients. Seed and seedlings can normally be protected when seed is treated at a rate of from 0.1 to 10 g of aggregate active ingredient per kilogram of seed.

The following Examples demonstrate the composition and method of the present invention and provide experimental evidence for synergy between the compound of Formula Ia and flusilazole in preventative control of wheat leaf rust caused by *Puccinia recondita*, wheat foot rot caused by *Pseudocercosporella herpotrichoides*, and wheat glume blotch caused by *Septoria nodorum*. The experimental also provides evidence for synergy between the compound of Formula Ia and tebuconazole in preventative control of wheat leaf rust and wheat glume blotch. The experimental also provides evidence for synergy between the compound of Formula Ia and flusilazole in curative control of wheat powdery mildew caused by *Erysiphe graminis f. sp. tritici* and wheat leaf rust.

The experimental also provides evidence for synergy between the compounds of Formula Ia, Formula IIa and flusilazole in preventative control of wheat foot rot. The experimental also provides evidence for synergy between the compounds of Formula Ia, IIa and flusilazole in curative control of wheat powdery mildew. The pathogen control protection afforded by these compositions is not limited, however, to these species.

The following TESTS demonstrate the control efficacy of compounds of this invention on specific pathogens. The pathogen control protection afforded by the compounds is not limited, however, to these species.

BIOLOGICAL EXAMPLES OF THE INVENTION

Test compounds were first dissolved in acetone in an amount equal to 50% of the final volume and then suspended at a concentrations from 0.08 to 200 ppm in purified water containing 250 ppm of the surfactant Trem[®] 014 (polyhydric alcohol esters). The resulting test suspensions were then used in the following test protocols. Spraying these test suspensions to the point of run-off on the test plants is the equivalent of a rate

Protocol #1 - WPM Preventive

The test compounds were sprayed to the point of run-off on wheat seedlings. The following day the seedlings were inoculated with a spore dust of *Erysiphe graminis* f. sp. tritici, (the causal agent of wheat powdery mildew). Seedlings were incubated in a growth chamber at 20 °C for 6 days, after which disease ratings were made.

Protocol #2 - WPM Curative

Wheat seedlings were inoculated with a spore dust of *Erysiphe graminis* f. sp. tritici, (the causal agent of wheat powdery mildew). The following day a test compounds were sprayed to the point of run-off on and seedlings incubated in a growthchamber at 20 °C for 7 days, after which disease ratings were made.

Protocol #3 WLR Preventive

The test compounds were sprayed to the point of run-off on wheat seedlings. The following day the seedlings were inoculated with a spore suspension of *Puccinia recondita* (the causal agent of wheat leaf rust) and incubated in a saturated atmosphere at 20 °C for 24 h, and then moved to a growth chamber at 20 °C for 6 days, after which disease ratings were made.

Protocol #4 - WLR Curative

Wheat seedlings were inoculated with a spore suspension of *Puccinia recondita* (the causal agent of wheat leaf rust). The following day the test compounds were sprayed to the point of run-off on and incubated in a saturated atmosphere at 20 °C for 24 h, then moved to a growth chamber at 20 °C for 6 days, after which disease ratings were made.

Protocol #5 - WFR Preventive

The test compounds were sprayed to the point of run-off on wheat seedlings. The following day the seedlings were inoculated with a spore suspension of *Pseudocercosporella herpotrichoides* (the causal agent of wheat eye spot or wheat foot rot) and incubated in a saturated atmosphere at 20 °C for 72 h, and then moved to a growth chamber at 20 °C for 6 days, after which disease ratings were made.

Protocol #6 - WGB Preventive

The test compounds were sprayed to the point of run-off on wheat seedlings. The following day the seedlings were inoculated with a spore suspension of *Septoria*

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of 500 g/ha.

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nodorum (the causal agent of wheat glume blotch) and incubated in a saturated atmosphere at 20 °C for 48 h, and then moved to a growth chamber at 20 °C for 6 days, after which disease ratings were made.

Protocol #7 - WGB Curative

Wheat seedlings were inoculated with a spore suspension Septoria nodorum (the causal agent of wheat glume blotch). Two days later test compounds were sprayed to the point of run-off on and seedlings incubated in a saturated atmosphere at 20 °C for 48 h, then moved to a growth chamber at 20 °C for 6 days, after which disease ratings were made.

Protocol #8 - WPM Preventative

The test suspension was sprayed to the point of run-off on wheat seedlings. The following day the seedlings were inoculated with a spore dust of *Erysiphe graminis f. sp. tritici*, (the causal agent of wheat powdery mildew) and incubated in a growth chamber at 20 °C for 7 days, after which disease ratings were made.

Results for protocols 1 - 8 are given in Tables A-G. In the tables, a rating of 100 indicates 100% disease control and a rating of 0 indicates no disease control (relative to the controls). An (nt) indicates no test results and (--) indicates no estimate of activity. In Tables A-G rates are given in parts per million (ppm) and/or grams active ingredient per hectare (g a.i./ha). The term "Actual" stands for the experimental value. The abbreviation "Exp." stands for "Expected" (i.e., the predicted activity, p, from the Colby equation).

TABLE A

Synergistic Effects of Compound Ia in Combination with Flusilazole, Tebuconazole and Prochloraz for Preventive Control of

	$\overline{\text{WGB}^{(4)}}$	ual Exp.	1	;	i	;	1	;	;	!	;	0	0	0	0	09	83	100
		Actual	0	0	0	0	0	0	0	09	83	100	0	0	0	43	91	66
<u>lol</u>	WFR ⁽³⁾	Exp.	1	i	1	ł	ł	1	i	1	ŀ	!	16	54	45	50	100	100
sease Contro	WF	<u>Actual</u>	0	0	0	0	16	54	45	50	100	100	0	∞	100	81	100	100
Percent Disease	WLR ⁽²⁾	Exp.	ł	ţ	ł	1	ŀ	:	ŀ	i	;	;	18	17	88	27	79	100
ă.	M	Actual	0	6	0	0	18	6	88	27	79	100	6	18	88	32	86	66
	M ₍₁₎	Exp. (6)	(2)	1	1	;	1	1	1	ł	1	\$ \$	100	100	100	100	ł	ŀ
	WPM ⁽¹⁾	Actual ⁽⁵⁾	100	100	100	100	06	66	100	62	35	81	100	100	100	100	nt(8)	nt
	Mole	Ratio											1.18:1	1.18:1	1.18:1	1:1.69	2.95:1	11.81:1
	Rates	(g a.i./ha)	0.2		5	25	0.2		Ś	2.5	12.5	50	0.2 + 0.2	1+1	5+5	2.5 + 5	12.5 + 5	50 + 5
	Rates	(mdd)	80.0	0.4	2	10	80.0	0.4	2	_	Ş	20	0.08 + 0.08 0.2	0.4 + 0.4	2 + 2	1+2	5+2	20 + 2
	Test	Cmpd 2	!	;	;	;	;	;	1	1	;	;	Ia	. Ia	Ia	Ia	la	ľa
	Test	Cmpd 1	Ia	la	Ia	Ia	Flusilazole	Flusilazole	Flusilazole	Flusilazole	Flusilazole	Flusilazole						

	€ I	Exp.	09	83	100	ł	;	:	;	:	55	96	100	:	55	96	100	;	:	}	;	;
	WGB ⁽⁴⁾	Actual	38	94	100	nt	55	96	100	{	59	100	100	nt	84	100	100	nt	nt	nt	nt	nt
731	6]	Exp.	50	100	100	;	ł	;	ł	;	;	:	ł	;	;	ł	ł	ł	í	1	37	100
Percent Disease Control	WFR ⁽³⁾	Actual	96	100	100	nt	37	100	100	37	100											
rcent Dise	R ⁽²⁾	Exp.	27	79	100	ł	ł	ŀ	ŀ	32	100	100	;	32	100	100	ł	ŀ	ŀ	ł	ł	i
al al	$\overline{\text{WLR}^{(2)}}$	Actual	70	100	100	32	100	100	nt	86	100	100		9/	66	100	nt	nt	nt	nt	nt	nt
	(()	Exp. (6)	ł	ŧ	;	1	;	{	1	1	;	1	i	1	1	1	1	1	1	;	1	;
	WPM ⁽¹⁾	Actual ⁽⁵⁾	nt	nt	nt	nt	nt	nt	nt	nt	nt	nt	nt	nt	nt	nt	nt	nt	nt	nt	nt	nt
	Mole	Ratio	1:8.47	1:1.69	2.36:1					1:4.13	1:1.21	6.06:1	24.23:1	1:20.63	1:4.13	1:1.21	4.85:1				1:5	1:1
	Rates	(g a.i./ha)	2.5 + 25	12.5 + 25	50 + 25	1	\$	25	100	1 + 5	5+5	25 + 5	100 + 5	1 + 25	5+25	25 + 25	100 + 25	-	5	25	1+5	5+5
	Rates			5 + 10					40				40 + 2									2 + 2
	Test	Cmpd 2	Ia	Га	la	ł	I E		ł	Ia	Ia	Ia	ľa	Ia	Ia	Ia	Ia	{	i	1	Ia	la
	Test	Cmpd 1	Flusilazole	Flusilazole	Flusilazole	Tebuconazole	Prochloraz	Prochloraz	Prochloraz	Prochloraz	Prochloraz											

	E	Exp.	;	;	:	;
				nt		
[o]	©)	Exp.	100	37	100	100
sease Contr	$\overline{\text{WLR}^{(2)}}$ $\overline{\text{WFR}^{(3)}}$	Actual	100	37	100	100
ercent Dis	R ⁽²⁾	Exp.	1	1	;	:
Pe	MI	Actual	nt	nt	nt	nt
	M()	Exp. (6)	1	;	ļ	:
	WPI	Actual ⁽⁵⁾	nt	nt	nt	Ħ
	Mole	Ratio	5:1	1:25	1:5	1::
	Rates	(g a.i./ha)	25 + 5	1 + 25	5 + 25	25 + 25
	Rates	(mdd)	10 + 2	0.4 + 10	2 + 10	10 + 10
	Test	Cmpd 2	la	Ia	Ia	Ia
	Test	Cmpd 1	Prochloraz	Prochloraz	Prochloraz	Prochloraz

(1) Wheat powdery mildew activity - Protocol #1

(2) Wheat leaf rust activity - Protocol #3

(3) Wheat foot rot activity - Protocol # 5

(4) Wheat glume blotch activity - Protocol # 6

(5) Actual activity observed in the test

(6) Predicted activity

(7) No estimate - (--) (8) Not tested (nt)

w (WPM), TABLEB

erv Mildew			WGB ⁽³⁾	Exp.	;	;	;	ł	ł	;	;	;	;	;	:	:	ì	15	23	72
/heat Powd		T-01	W	Actual	0	0	10	0	0	0	5	0	15	69	nţ	nt	nt	0	43	71
ntrol of W		ease Contro	$WLR^{(2)}$	Exp.	;	:	;	1	1	1	;		:	;	1	;	;	43	84	100
r Chrative Co	Wheat Leaf Rust (WLR) and Wheat Glume Blotch (WGB)	Percent Disease Control	WL	<u>Actual</u>	0	0	0	0	0	0	43	80	84	100	nt	nt	nt	15	100	66
onazole fo	Glume Bl		WPM ⁽¹⁾	Exp.	;	:	:	1	1	1	;	1	1	1	89	82	100	ţ	;	;
1ABLE B	and Wheat		WP	Actual	0.	99	09	nt	89	73	89	100	91	100	86	100	100	nt	nt	nt
ith Elucilazo	Rust (WLR)		Mole	Ratio											1.18:1	1.18:1	1.18:1	1:1.69	2.95:1	11.81:1
ombination w	Wheat Leaf		Rates	(g a.i./ha)	0.2		'n	25	0.2	-	2.5	Š	12.5	20	0.2 + 0.2	1+1	5+5	2.5 + 5	12.5 + 5	50 + 5
Out of Is in C			Rates	(mdd)	0.08	0.4	C1	10	0.08	0.4	1	C 1	5	20	0.08 + 0.08	0.4 + 0.4	2 + 2	1+2	5+2	20 + 2
Foots of Com			Test	Cmpd 2	;	1	;	;	ŀ	;	1	1	;		Ia	Ia	la	Ia	Ia	la
<u>1ABLE B</u> Semandistic Effects of Compound 1s in Combination with Flusilazole or Tehnonazole for Curative Control of Wheat Powdery Mildew 1			Test	Cmpd 1	Ia	la	Ia	Ia	Flusilazole											

	B ⁽³⁾	Exp.	0	15	69	ŀ	;	;	1	1	49	55	91	:	43	50	06
	WGB ⁽³⁾	Actual	0	29	81	nt	43	50	06	nt	29	71	90	nt	10	71	100
ease Contro	R ⁽²⁾	Exp.	43	84	100	:	;	;	;	40	100	100	100	40	100	100	100
Percent Disease Contro	WLR ⁽²⁾	Actual	71	100	100	40	100	100	100	100	100	100	100	100	100	100	100
	M ⁽¹⁾	Exp.	;	1	1	ł	1	;	;	;	;	ŧ	;	1	1	1	!
	WPM ⁽¹⁾	<u>Actual</u>	nt	nt	nt	nt	nt	nt	nt	nt	nt	nt	nt	nt	nt	nt	nt
	Mole	Ratio	1:8.47	1:1.69	2.36:1					1:4.13	1:1.21	6.06:1	24.23:1	1:20.63	1:4.13	1:1.21	4.85:1
	Rates	(g a.i./ha)	2.5 +25	12.5 + 25	50 + 25	_	2	25	100	1 + 5	5+5	25 + 5	100 + 5	1 + 25	5 + 25	25 + 25	100 + 25
	Rates	(mdd)	1 + 10	5 + 10	20 + 10	0.4	2	10	40	0.4 + 2	2+2	10 + 2	40 + 2	0.4 + 10	2 + 10	10 + 10	40 + 10
	Test	Cmpd 2	Ia	Ia	Ia	:	;	;		Ia	Ia	Ia	Ia	la	Ia	Ia	<u>e</u>
	Test	Cmpd 1	Flusilazole	Flusilazole	Flusilazole	Tebuconazole											

(1) Wheat powdery mildew - Protocol # 2

⁽²⁾ Wheat leaf rust - Protocol # 4(3) Wheat glume blotch - Protocol # 7

Synergistic Effects of Compound Ia in Combination with Compound IIa for Preventive Control of Wheat Powdery Mildew (WPM), TABLEC

Wheat Leaf Rust (WLR), Wheat Foot Rot (WFR) and Wheat Glume Blotch (WGB)

							٦	- 2		
B(±)	Exp.	;	ł	;	:	1	!	0	0	0
MG	Actual	0	0	0	0	0	0	0	0	0
R ⁽³⁾	Exp.	ł	1	1	1	!	ł	∞	26	38
WF	Actual	0	0	0	. 8	26	38	0	∞	61
R ⁽²⁾	Exp.	ł	:	ŀ	;	:	:	54	92	91
WE	Actual	0	6	0	54	91	66	54	93	66
() ()	Exp.	}	ŧ	:	1	ŧ	;	100	100	100
WP	Actual	100	100	100	09	88	100	66	100	100
Mole	Ratio							1:1.13	1:1.13	1:1.13
Rates	(g a.i./ha)	0.2	1	5	0.2	-	S	0.2 + 0.2	1+1	5 + 5
Rates	(mdd)	80.0	0.4	2	0.08	0.4	7	0.08 + 0.08	0.4 + 0.4	2+2
Test	Cmpd 3	ł	;	ł	ł	f	;	{	1	i
Test	Cmpd 2	1	1	;	1	1	;	Ia	Ia	ľa
Test	Cmpd 1	Ia	la	Ia	IIa	IIa	IIa	IIa	IIa	IIa

(3) Wheat foot rot activity - Protocol # 5 (2) Wheat leaf rust activity - Protocol # 3

(1) Wheat powdery mildew activity - Protocol #1

(4) Wheat glume blotch activity - Protocol # 6

TABLED

Synergistic Effects of Compound Ia in Combination with Compound IIa on Curative Control of Wheat Powdery Mildew (WPM), Wheat Leaf Rust (WLR) and Wheat Glume Blotch (WGB)

	B ⁽³⁾	Exp.	;	1	;	ţ	ŀ	ŀ	13	0	22
힏	WG	Actual	0	- 0	10	13	0	13	0	13	53
ise Conti		Exp.	:	ł	;	ŀ	:	!	0	0	0
rcent Disease				- 0							0
Pe	WPM(1)	Exp.		1	ì	;	ļ	ł	0	88	100
	WPI	Actual	0	95	09	0	73	100	21	66	100
	Mole	Ratio							1:1.13	1:1.13	1:1.13
	Rates	(g a.i./ha)	0.2	•	5	0.2		5	0.2 + 0.2	1+1	5 + 5
	Rates	(mdd)	0.08	0.4	7	0.08	0.4	2	0.08 + 0.08	0.4 + 0.4	2 + 2
	Test	Cmpd 3	ł	ŀ	ŀ	:	;	ŀ	;	ļ	;
	Test	Cmpd 2		;	:	:	;	;	Ia	Ia	Ia
	Test	Cmpd 1	la	la	Ia	IIa	IIa	IIa	IIa	Ila	IIa

(1) Wheat powdery mildew - Protocol # 2

⁽²⁾ Wheat leaf rust - Protocol # 4

⁽³⁾ Wheat glume blotch - Protocol # 7

Synergistic Effects of Compound Ia / Compound IIa / Flusilazole combinations for Preventive Control of Wheat Powdery Mildew (WPM), Wheat Leaf Rust (WLR), Wheat Foot Rot (WFR) and Wheat Glume Blotch (WGB) TABLEE

							44	Ŧ						
	B ⁽⁴⁾	Exp.	;	ì	;	:	;	;	ł	1	ł	0	0	0
	WGB(4)	Actual	0	0	0	0	0	0	0	0	0	0	0	47
trol	(£)	Exp.	;	;					;				54	45
Percent Disease Control	WFR ⁽³⁾	Actual	0	0	0	8	26	38	16	54	45	∞	62	100
ent Dis	WLR ⁽²⁾	Exp.	1	ł	1				i		;	62	100	100
Perc	WL	Actual	0	6	0	54	16	66	18	6	88	28	93	66
	()	Exp.	i	1	1				1		!	100	100	100
	WPM(1)	Actual	100	001	100	09	88	100	06	66	100	100	100	100
	Mole	Ratio										1.33:1.13:1	1.33:1.13:1	1.33:1.13:1
		_		1							5	80.2 + 0.2 + 0.2	1 + 1 + 1	5 + 5 + 5
	Rates	(mdd)	0.08	0.4	7	80.0	0.4	C 1	80.0	0.4	2	la $0.08 + 0.08 + 0.08$ (0.4 + 0.4 + 0.4	2+2+2
	Test	Cmpd 3	ŀ	;	1	!	;	i	ì	1	1	Ha	IIa	IIa
	Test	Cmpd 2	ł	4	;	ţ	ł	t	i	1	{	Ia	Ia	la
	Test	Cmpd 1	Ia	la	ľa	Ifa	Ha	IIa	Flusilazole	Flusilazole	Flusilazole	Flusilazole	Flusilazole	Flusilazole

⁽¹⁾ Wheat powdery mildew activity - Protocol #1

(3) Wheat foot rot activity - Protocol # 5

⁽⁴⁾ Wheat glume blotch activity - Protocol # 6

⁽²⁾ Wheat leaf rust activity - Protocol # 3

TABLEF

Synergistic Effects of Compound Ia / Compound IIa / Flusilazole on Curative Control of Wheat Powdery Mildew (WPM),

Wheat Leaf Rust (WLR) and Wheat Glume Blotch (WGB)

	36	Exp.	1	;	;	;		;	;	;	;	13	0	22
1	WGB ⁽⁵⁾	Actual Exp	0	0	10	13	0	13	0	0	0	40	0	13
e Contro	(3)	Exp.	;	i	;	ŀ	ł	;	1	ł	ł	0	0	80
Percent Disease Control	WLR ⁽²⁾	Actual	0	0	0	0	0	0	0	0	80	0	0	98
Perc	Ma	Exp.	1	ŀ	ł	;	1	1	1	;	1	89	100	100
	WPM ⁽¹⁾	Actual	0	99	09	0	73	100	89	73	100	96	100	100
	Mole	Ratio										1.33:1.13:1	1.33:1.13:1	1.33:1.13:1
	Rates	(g a.i./ha)	0.2		'n	0.2		Š	0.2	1	5	0.2 + 0.2 + 0.2	1+1+1	5+5+5
	Rates	(mdd)	0.08	0.4	61	80.0	0.4	C1	80.0	0.4	2	0.08 + 0.08 + 0.08	0.4 + 0.4 + 0.4	2 + 2 + 2
	Test	Cmpd 3	;	;	;	1	1	;	1	;	;	IIa	IIa	IIa
	Test	Cmpd 2	ţ	· { ·	;	;	;	£	;	;	;	Ia	Га	Ia
	Test	Cmpd 1	la	Га	la	Па	IIa	Ila	Flusilazole	Flusilazole	Flusilazole	Flusilazole	Flusilazole	Flusilazole

(1) Wheat powdery mildew - Protocol # 2

(2) Wheat leaf rust - Protocol # 4

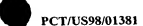
(3) Wheat glume blotch - Protocol # 7

(5) Not tested (nt)

TABLEG

Synergistic Effects of Compound Ia in Combination with Fenpropimorph for Preventative Control of Wheat Powdery Mildew (WPM)(1)

	Exp.(3)		(4)	:	;	:	ŀ	;	100	9.7.6	72					
	Mean		100	96	72	78	41	0	100	66	87				the test	
7PM(2)	Test 2 Test 3		100	64	34	80	13	0	100	86	82	52			erved in	
rol of W	Test 2		100	95	87	9/	92	nt(5)	100	100	nt	31			vity obs	() - e
% Control of WPM(2)	Test 1		100	96	94	6/	34	0	100	66	92	19			(2) Actual activity observed in the test	(4) No estimate - ()
	Mole	Ratio							6.1:1	4.9:1	6.1:1				(2) Ac	(4) No
	Rates (g a.i./ha)		20	· ·		100	20	5	100 + 20	20 + 5	5 + 1				- Protocol #8	
	Cmpd 2		;	;	;	;	ł	1	Ia	la	Ia				ildew activity	
	Cmpd 1		la	la	ľa	Fenpropimorph	Fenpropimorph	Fenpropimorph	Fenpropimorph	Fenpropimorph	Fenpropimorph	Untreated	(% area diseased	Foliage)	(1) Wheat powdery mildew activity - Protocol #8	(3) Predicted activity



CLAIMS

What is claimed is:

- 1. A fungicidal composition comprising:
- (1) at least one compound selected from the quinazolinones of Formula I,
- 5 N-oxides, and agriculturally suitable salts thereof,

wherein

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D is O or S;

10 R^1 is C_1 - C_{10} alkyl; C_3 - C_5 cycloalkyl; C_3 - C_{10} alkenyl; C_3 - C_{10} alkynyl;

C₁-C₁₀ haloalkył; C₃-C₁₀ haloalkenyl; C₃-C₁₀ haloalkynyl;

C₂-C₁₀ alkoxyalkyl; C₂-C₁₀ alkylthioalkyl; C₂-C₁₀ alkylsulfinylalkyl;

C₂-C₁₀ alkylsulfonylalkyl; C₄-C₁₀ cycloalkylalkyl;

 C_4 - C_{10} alkenyloxyalkyl; C_4 - C_{10} alkynyloxyalkyl; C_4 - C_{10}

(cycloalkyl)oxyalkyl; C₄-C₁₀ alkenylthioalkyl; C₄-C₁₀ alkynylthioalkyl;

C₄-C₁₀ (cycloalkyl)thioalkyl; C₂-C₁₀ haloalkoxyalkyl;

C₄-C₁₀ haloalkenyloxyalkyl; C₄-C₁₀ haloalkynyloxyalkyl;

 C_4 - C_{10} alkoxyalkenyl; C_4 - C_{10} alkoxyalkynyl; C_4 - C_{10} alkylthioalkenyl;

C₄-C₁₀ alkylthioalkynyl; C₄-C₁₀ trialkylsilylalkyl; C₁-C₁₀ alkyl substituted

with NR⁵R⁶, nitro, cyano or phenyl optionally substituted with R⁸, R⁹ and

 $R^{10};\,C_1\text{-}C_{10}$ alkoxy; $C_1\text{-}C_{10}$ haloalkoxy; $C_1\text{-}C_{10}$ alkylthio;

C₁-C₁₀ haloalkylthio; or pyridinyl, furanyl, thienyl, naphthalenyl,

benzofuranyl, benzothienyl or quinolinyl each optionally substituted with

R8, R9 and R10:

R² is C_1 - C_{10} alkyl; C_3 - C_7 cycloalkyl; C_3 - C_{10} alkenyl; C_3 - C_{10} alkynyl;

C₁-C₁₀ haloalkyl; C₃-C₁₀ haloalkenyl; C₃-C₁₀ haloalkynyl;

C₂-C₁₀ alkoxyalkyl; C₂-C₁₀ alkylthioalkyl; C₂-C₁₀ alkylsulfinylalkyl;

C₂-C₁₀ alkylsulfonylalkyl; C₄-C₁₀ cycloalkylalkyl;

C₄-C₁₀ alkenyloxyalkyl; C₄-C₁₀ alkynyloxyalkyl; C₄-C₁₀

30 (cycloalkyl)oxyalkyl; C₄-C₁₀ alkenylthioalkyl; C₄-C₁₀ alkynylthioalkyl;

C₄-C₁₀ (cycloalkyl)thioalkyl; C₂-C₁₀ haloalkoxyalkyl;

C₄-C₁₀ haloalkenyloxyalkyl; C₄-C₁₀ haloalkynyloxyalkyl;

 C_4 - C_{10} alkoxyalkenyl; C_4 - C_{10} alkoxyalkynyl; C_4 - C_{10} alkylthioalkenyl;

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 C_4 - C_{10} alkylthioalkynyl; C_4 - C_{10} trialkylsilylalkyl; C_2 - C_{10} cyanoalkyl; C_1 - C_{10} nitroalkyl; C_1 - C_{10} alkyl substituted with CO_2R^5 , NR^5R^6 , or phenyl optionally substituted with R^7 , R^9 and R^{10} ; phenyl optionally substituted with R^7 , R^9 and R^{10} ; -N= $-CR^5R^5$; or $-NR^5R^6$; or

- 5 R¹ and R² are taken together to form -CH₂(CH₂)_qCH₂-; q is 0, 1, 2, 3 or 4;
 - R³ is halogen, C₁-C₈ alkyl, C₃-C₈ cycloalkyl, C₂-C₈ alkenyl, C₂-C₈ alkynyl, C₁-C₈ haloalkyl, C₃-C₈ haloalkenyl, C₃-C₈ haloalkynyl, C₁-C₈ alkoxy, C₁-C₈ haloalkoxy, C₃-C₈ alkenyloxy, C₃-C₈ alkynyloxy, C₁-C₈ alkylthio, C₃-C₈ alkenylthio, C₁-C₈ alkylsulfinyl, C₁-C₈ alkylsulfinyl, C₁-C₈ alkylsulfinyl, C₂-C₈ alkylsulfinylalkyl, C₂-C₈ alkylsulfinylalkyl, C₂-C₈ alkylsulfonylalkyl, C₄-C₈ cycloalkylalkyl, C₃-C₈ trialkylsilyl, NR⁵R⁶, C₅-C₈ trialkylsilylalkynyl or phenyl optionally substituted with at least one R⁷;
- 15 R⁴ is hydrogen, halogen, C₁-C₄ alkyl, C₁-C₄ haloalkyl, C₁-C₄ alkoxy or C₁-C₄ haloalkoxy;
 - each R⁵ is independently hydrogen, C₁-C₄ alkyl or phenyl optionally substituted with at least one R⁷;
 - each R^6 is independently hydrogen, C_1 - C_8 alkyl or phenyl optionally substituted with at least one R^7 ; or
 - when R^5 and R^6 are attached to the same nitrogen atom, R^5 and R^6 can be taken together to form -CH₂CH₂CH₂CH₂-, -CH₂(CII₂)₃CH₂-, -CH₂CH₂OCH₂CH₂-, -CH₂CH(Me)CH₂- or -CH₂CH(Me)OCH(Me)CH₂-;
 - each R⁷ is independently halogen, C₁-C₄ alkyl, C₁-C₄ alkoxy, C₁-C₄ haloalkyl, nitro or cyano;
 - R⁸ is C₁-C₆ alkyl, C₁-C₆ alkoxy, C₁-C₆ haloalkyl, halogen, C₂-C₈ alkynyl, C₁-C₆ alkylthio, phenyl or phenoxy each optionally substituted with at least one R⁷, cyano, nitro, C₁-C₆ haloalkoxy, C₁-C₆ haloalkylthio, C₂-C₆ alkenyl, C₂-C₆ haloalkenyl, acetyl, CO₂Me or N(C₁-C₂ alkyl)₂;
 - each R⁹ is independently methyl, ethyl, methoxy, methylthio, halogen or trifluoromethyl; and
 - each R¹⁰ is independently halogen;
- (2) at least one compound selected from (a) compounds acting at the *bc*₁ complex of the fungal mitochondrial respiratory electron transfer site and (b) compounds that control fungal disease by inhibiting sterol biosynthesis; and optionally

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- (3) at least one of a surfactant, a solid diluent or a liquid diluent; wherein component (1) and component (2) are present in a fungicidally effective amount and the mole ratio of component (1) to component (2) is from about 30:1 to 1:30.
- 2. The fungicidal composition of Claim 1 comprising (1) at least one compound selected from the group consisting of

6-bromo-3-propyl-2-propyloxy-4(3*H*)-quinazolinone,

6,8-diiodo-3-propyl-2-propyloxy-4(311)-quinazolinone and

6-iodo-3-propyl-2-propyloxy-4(3H)-quinazolinone;

propiconazole and tebuconazole.

and (2) at least one compound selected from the group consisting of

2,4-dihydro-5-methoxy-2-methyl-4-[2-[[[[1-[3-(trifluoromethyl)phenyl]ethylidene]amino]oxy]methyl]phenyl]-3*H*-1,2,4triazol-3-one, kresoxim-methyl, azoxystrobin, (*E*)-2-(methoxyimino)-*N*methyl-2-(2-phenoxyphenyl)acetamide, flusilazole, epoxiconazole,

3. The fungicidal composition of Claim 2 comprising a fungicidally effective amount of (1) 6-iodo-3-propyl-2-propyloxy-4(3*H*)-quinazolinone and (2) 2,4-dihydro-5-methoxy-2-methyl-4-[2-[[[1-[3-

(trifluoromethyl)phenyl]ethylidene]amino[oxy]methyl]phenyl]-3*H*-1,2,4-triazol-3-one.

- 4. The fungicidal composition of Claim 2 comprising a fungicidally effective amount of (1) 6-iodo-3-propyl-2-propyloxy-4(3*H*)-quinazolinone and (2) flusilazole.
- 5. The fungicidal composition of Claim 2 comprising a fungicidally effective amount of (1) 6-iodo-3-propyl-2-propyloxy-4(3*H*)-quinazolinone and (2) tebuconazole.
- 6. The fungicidal composition of Claim 2 comprising a fungicidally effective amount of (1) 6-iodo-3-propyl-2-propyloxy-4(3*H*)-quinazolinone and (2) both 2,4-dihydro-5-methoxy-2-methyl-4-[2-[[[[1-[3-(trifluoromethyl)phenyl]ethylidene]amino]oxy]methyl]phenyl]-3*H*-1,2,4-triazol-3-one; and flusilazole.
- 7. A method for controlling plant diseases caused by fungal plant pathogens comprising applying to the plant or portion thereof, or to the plant seed or seedling, a fungicidally effective amount of a composition of Claim 1.
- 8. A method of controlling at least one fungal plant disease selected from *Erysiphe graminis, Puccinia recondita, Pseudocercosporella herpotrichoides* and *Septoria nodorum* comprising applying to the plant or portion thereof, or to the plant seed or seedling
- 35 (1) at least one compound selected from the quinazolinones of Formula I, *N*-oxides, and agriculturally suitable salts thereof,

wherein

D is O or S;

5 R^1 is C_1 - C_{10} alkyl; C_3 - C_5 cycloalkyl; C_3 - C_{10} alkenyl; C_3 - C_{10} alkynyl; C₁-C₁₀ haloalkyl; C₃-C₁₀ haloalkenyl; C₃-C₁₀ haloalkynyl; C_2 - C_{10} alkoxyalkyl; C_2 - C_{10} alkylthioalkyl; C_2 - C_{10} alkylsulfinylalkyl; C₂-C₁₀ alkylsulfonylalkyl; C₄-C₁₀ cycloalkylalkyl; C₄-C₁₀ alkenyloxyalkyl; C₄-C₁₀ alkynyloxyalkyl; C₄-C₁₀ 10 (cycloalkyl)oxyalkyl; C₄-C₁₀ alkenylthioalkyl; C₄-C₁₀ alkynylthioalkyl; C₄-C₁₀ (cycloalkyl)thioalkyl; C₂-C₁₀ haloalkoxyalkyl; C₄-C₁₀ haloalkenyloxyalkyl; C₄-C₁₀ haloalkynyloxyalkyl; C_4 - C_{10} alkoxyalkenyl; C_4 - C_{10} alkoxyalkynyl; C_4 - C_{10} alkylthioalkenyl; C₄-C₁₀ alkylthioalkynyl; C₄-C₁₀ trialkylsilylalkyl; C₁-C₁₀ alkyl substituted with NR⁵R⁶, nitro, cyano or phenyl optionally substituted with R⁸, R⁹ and 15 R^{10} ; C_1 - C_{10} alkoxy; C_1 - C_{10} haloalkoxy; C_1 - C_{10} alkylthio; C₁-C₁₀ haloalkylthio; or pyridinyl, furanyl, thienyl, naphthalenyl, benzofuranyl, benzothienyl or quinolinyl each optionally substituted with R8, R9 and R10; 20 R^2 is C_1 - C_{10} alkyl; C_3 - C_7 cycloalkyl; C_3 - C_{10} alkenyl; C_3 - C_{10} alkynyl; C₁-C₁₀ haloalkyl; C₃-C₁₀ haloalkenyl; C₃-C₁₀ haloalkynyl; C₂-C₁₀ alkoxyalkyl; C₂-C₁₀ alkylthioalkyl; C₂-C₁₀ alkylsulfinylalkyl; C₂-C₁₀ alkylsulfonylalkyl; C₄-C₁₀ cycloalkylalkyl; C₄-C₁₀ alkenyloxyalkyl; C₄-C₁₀ alkynyloxyalkyl; C₄-C₁₀ 25 (cycloalkyl)oxyalkyl; C₄-C₁₀ alkenylthioalkyl; C₄-C₁₀ alkynylthioalkyl; C₄-C₁₀ (cycloalkyl)thioalkyl; C₂-C₁₀ haloalkoxyalkyl; C₄-C₁₀ haloalkenyloxyalkyl; C₄-C₁₀ haloalkynyloxyalkyl; C_4 - C_{10} alkoxyalkenyl; C_4 - C_{10} alkoxyalkynyl; C_4 - C_{10} alkylthioalkenyl; C_4 - C_{10} alkylthioalkynyl; C_4 - C_{10} trialkylsilylalkyl; C_2 - C_{10} cyanoalkyl; C_1 - C_{10} nitroalkyl; C_1 - C_{10} alkyl substituted with CO_2R^5 , NR^5R^6 , or phenyl 30 optionally substituted with R⁷, R⁹ and R¹⁰; phenyl optionally substituted with R^7 , R^9 and R^{10} ; -N=CR⁵R⁵; or -NR⁵R⁶; or

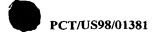
 R^1 and R^2 are taken together to form -CH₂(CH₂)_qCH₂-;

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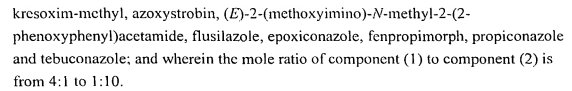




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q is 0, 1, 2, 3 or 4;

- R³ is halogen, C¹-C² alkyl, C³-C² cycloalkyl, C²-C² alkenyl, C²-C² alkynyl, C¹-C² haloalkyl, C³-C² haloalkenyl, C³-C² haloalkynyl, C¹-C² alkoxy, C¹-C² haloalkoxy, C³-C² alkenyloxy, C³-C² alkynyloxy, C¹-C² alkylthio, C³-C² alkenylthio, C³-C² alkynylthio, C¹-C² alkylsulfinyl, C¹-C² alkylsulfinyl, C²-C² alkylsulfinyl, C²-C² alkylsulfinylalkyl, C²-C² alkylsulfinylalkyl, C²-C² alkylsulfinylalkyl, C³-C² alkylsulfinylalkyl, C³-C² trialkylsilyl, NR⁵R², C³-C² trialkylsilylalkynyl or phenyl optionally substituted with at least one R³;
- 10 R⁴ is hydrogen, halogen, C₁-C₄ alkyl, C₁-C₄ haloalkyl, C₁-C₄ alkoxy or C₁-C₄ haloalkoxy;
 - each R^5 is independently hydrogen, C_1 - C_4 alkyl or phenyl optionally substituted with at least one R^7 ;
 - each R⁶ is independently hydrogen, C₁-C₈ alkyl or phenyl optionally substituted with at least one R⁷; or
 - when R⁵ and R⁶ are attached to the same nitrogen atom, R⁵ and R⁶ can be taken together to form -CH₂CH₂CH₂CH₂-, -CH₂(CH₂)₃CH₂-, -CH₂CH₂OCH₂CH₂-, -CH₂CH(Me)CH₂- or -CH₂CH(Me)OCH(Me)CH₂-;
 - each R⁷ is independently halogen, C₁-C₄ alkyl, C₁-C₄ alkoxy, C₁-C₄ haloalkyl, nitro or cyano;
 - R^8 is C_1 - C_6 alkyl, C_1 - C_6 alkoxy, C_1 - C_6 haloalkyl, halogen, C_2 - C_8 alkynyl, C_1 - C_6 alkylthio, phenyl or phenoxy each optionally substituted with at least one R^7 , cyano, nitro, C_1 - C_6 haloalkoxy, C_1 - C_6 haloalkylthio, C_2 - C_6 alkenyl, C_2 - C_6 haloalkenyl, acetyl, CO_2 Me or $N(C_1$ - C_2 alkyl)₂;
 - each R⁹ is independently methyl, ethyl, methoxy, methylthio, halogen or trifluoromethyl; and
 - each R10 is independently halogen; and
- (2) at least one compound selected from (a) compounds acting at the bc_1 complex of the fungal mitochondrial respiratory electron transfer site and (b) compounds that control fungal disease by inhibiting sterol biosynthesis; wherein component (1) and component (2) are added in amounts sufficient to provide a fungicidal effectiveness greater than the sum of the fungicidal effectivenesses provided by those amounts of said components taken independently.
- 9. A method of Claim 8 wherein *Erysiphe graminis* is controlled; wherein component (1) is 6-iodo-3-propyl-2-propyloxy-4(3*H*)-quinazolinone and component (2) is selected from the group consisting of 2,4-dihydro-5-methoxy-2-methyl-4-[2-[[[1-[3-(trifluoromethyl)phenyl]ethylidene]amino]oxy]methyl]phenyl]-3*H*-1,2,4-triazol-3-one,



- 5 10. A method of Claim 8 wherein at least one of *Puccinia recondita* and *Septoria nodorum* is controlled; wherein component (1) is 6-iodo-3-propyl-2-propyloxy-4(3*H*)-quinazolinone and component (2) is selected from the group consisting of 2,4-dihydro-5-methoxy-2-methyl-4-[2-[[[[1-[3-(trifluoromethyl)phenyl]ethylidene]amino]oxy]methyl]phenyl]-3*H*-1,2,4-triazol-3-one, kresoxim-methyl, azoxystrobin, (*E*)-2-(methoxyimino)-*N*-methyl-2-(2-phenoxyphenyl)acetamide, flusilazole, epoxiconazole, fenpropimorph, propiconazole and tebuconazole; and wherein the mole ratio of component (1) to component (2) is from 30:1 to 1:30.
- 11. A method of Claim 8 wherein *Pseudocercosporella herpotrichoides* is controlled; wherein component (1) is 6-iodo-3-propyl-2-propyloxy-4(3*H*)-quinazolinone and component (2) is selected from the group consisting of 2,4-dihydro-5-methoxy-2-methyl-4-[2-[[[[1-[3-(trifluoromethyl)phenyl]ethylidene]amino]oxy]methyl]phenyl]-3*H*-1,2,4-triazol-3-one, kresoxim-methyl, azoxystrobin, (*E*)-2-(methoxyimino)-*N*-methyl-2-(2-phenoxyphenyl)acetamide, flusilazole, epoxiconazole, fenpropimorph, propiconazole and tebuconazole; and wherein the mole ratio of component (1) to component (2) is from 15:1 to 1:15.

INTERNATIONAL SEARCH REPORT

Int. .tiopal Application No

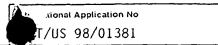
			PCT/Ú 8/01381	
A. CLASSI IPC 6	FICATION OF SUBJECT MATTER A01N43/54 //(A01N43/54,61:00,55 37:50)	5:00,47:38,43:	84,43:653,43:54,	
According to	o international Patent Classification(IPC) or to both national classifica	ation and IPC		
	SEARCHED			
IPC 6	ocumentation searched (classification system followed by classification AOIN	n symbols)		
Documental	tion searched other than minimum documentation to the extent that su	uch documents are include	d in the fields searched	
Electronic d	ata base consulted during the international search (name of data bas	se and, where practical, s	earch terms used)	
C. DOCUM	ENTS CONSIDERED TO BE RELEVANT			
Category °	Citation of document, with indication, where appropriate, of the rele	evant passages	Relevant to claim No.	
X	WO 94 26722 A (DU PONT ;BEREZNAK FRANCIS (US); CHANG ZEN YU (US); THO) 24 November 1994 cited in the application see page 1, line 10 - page 4, lir see page 9, line 18 - line 23	SELBY	1,2,4,5,	
Y	see page 33, line 3 - line 8 see page 33, line 16 - line 27 see page 33, line 33 - line 35	-/	2,6	
X Furt	her documents are listed in the continuation of box C.	X Patent family mo	embers are listed in annex.	
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